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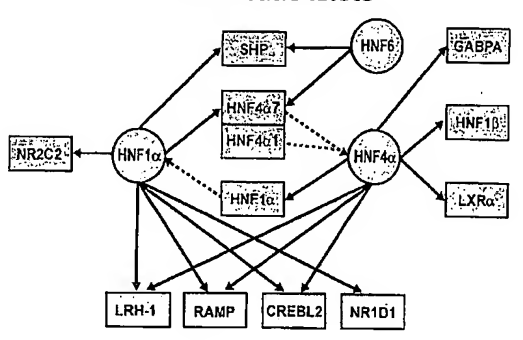
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(54) Title: TRANSCRIPTIONAL REGULATORS AND METHODS THEREOF

HNF1a HNF1a

(57) Abstract: The invention relates to transcriptional regulators and related methods thereof. The invention further relates to the identification of genes regulated by transcriptional regulators, to the treatment of diseases associated with abnormal function of a transcriptional regulator and to the modulation of gene expression, including genes expressed in hepatocytes or pancreatic cells, through the modulation of transcriptional regulator activity.

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Transcriptional Regulators and Methods Thereof

CROSS-REFERENCE TO RELATED APPLICATIONS

This application claims the benefit of the filing date of U.S. Application No. 60/525318, filed November 26, 2003, entitled "CONTROL OF PANCREAS AND LIVER GENE EXPRESSION BY HNF TRANSCRIPTION FACTORS", U.S. Application No. 60/542520, filed February 6, 2004, entitled "CONTROL OF PANCREAS AND LIVER GENE EXPRESSION BY HNF TRANSCRIPTION FACTORS", U.S. Application No. 60/544835, filed February 13, 2004, entitled "CONTROL OF PANCREAS AND LIVER GENE EXPRESSION BY HNF TRANSCRIPTION FACTORS", and U.S. Application No. 60/547933, filed February 26, 2004, entitled "TRANSCRIPTIONAL REGULATORS AND METHODS THEREOF". The entire teachings of the referenced applications are incorporated by reference herein.

FUNDING

The invention described herein was supported, in whole or in part, by the U.S. Department of Energy Program for Computational Molecular Biology. The United States government has certain rights in the invention.

BACKGROUND OF THE INVENTION

Gene expression is controlled by transcriptional regulatory proteins, which bind specific DNA sequences and recruit cofactors and the transcription apparatus to promoters (1-3). The expression of transcriptional regulators themselves is also regulated by transcriptional regulators, and a single gene may be regulated by multiple transcription factors. As a result of these regulatory networks, or pathways, misregulation of a single transcriptional regulator in a cell can result in the aberrant expression of multiple genes in the network in which the transcriptional regulator is active, leading to disease in the organism.

Current methods of identifying the genes controlled by a transcriptional regulator typically include a comparison of the mRNA levels of candidate target in

cells which express the transcriptional regulator and control cells which either do not express it. Often, this involves overexpressing a recombinant transcriptional regulator in a given cell type and using, as a control cell, one which overexpresses a control recombinant protein or no recombinant protein at all. However, given to the artificial nature of using cell lines and overexpressing transgenes, the results obtained from such approaches may not reflect the *in vivo* regulation by native transcriptional regulators in an organism.

Genome-wide analysis methods have been used recently to determine how tagged transcriptional regulators encoded in *Saccharomyces cerevisae* are associated with the genome in living yeast cells and to model the transcriptional regulatory circuitry of these cells (4). These methods have also been used in human tissue culture cells to identify target genes for several transcriptional regulators (5-7).

However, the need remains to develop genome-scale analysis methods to determine how transcriptional regulators control the global gene expression programs that characterize specific tissues, and in particular, freshly isolated, primary tissues, in which the transcriptional regulators are likely to maintain their *in vivo* specificities. Furthermore, there is a need to identify the regulatory networks or pathways in which a given transcriptional activator acts, in part, to allow for the identification of therapeutic targets for diseases caused by aberrant function of a transcriptional regulator.

SUMMARY OF THE INVENTION

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In one aspect, the invention provides a method of identifying the genes regulated by a transcriptional regulator. One aspect of the invention provides a method of determining which genes from a subset of genes are regulated by a transcriptional regulator in a cell, the method comprising (a) selectively isolating chromatin from a cell which expresses the transcriptional regulator to generate isolated chromatin; (b) selectively isolating chromatin fragments from the isolated chromatin to generate bound chromatin fragments, wherein the bound chromatin fragments are bound by the transcriptional regulator; (c) amplifying both the bound chromatin fragments to generate amplified chromatin fragments and the isolated chromatin to generate

amplified control chromatin; (d) hybridizing the amplified control chromatin and the amplified chromatin fragments to a DNA microarray, wherein the DNA microarray comprises (1) at least 10,000 experimental spots, each experimental spot comprising an experimental DNA, each experimental DNA comprising a promoter region from a gene in the subset; and (2) at least 100 control spots, each control spot comprising a control DNA, each control DNA comprising a non-promoter region; and (e) determining and comparing a hybridization signal at each of the spots on the microarray between those generated by (1) the amplified control chromatin; and (2) the amplified chromatin fragments; wherein a gene in the subset is said to be regulated by the transcriptional regulator in the cell if a spot comprising a promoter region of said gene displays a higher level of hybridization by the amplified chromatin fragments than by the amplified control chromatin.

In another aspect, the invention provides methods of identifying regulatory networks, or pathways, in a cell. The invention provides a method of identifying a transcriptional regulatory network in a cell, the method comprising determining if a transcriptional regulator regulates additional transcriptional regulators in the cell using the method of any of the methods described herein, wherein a transcriptional regulatory network is identified if at least one additional transcriptional regulator is regulated by the transcriptional regulator.

The invention also provides a method of identifying a transcriptional regulatory network in a cell, the method comprising determining if a transcriptional regulator regulates (i) its own promoter; or (ii) a promoter from a plurality of transcriptional regulators; using any of the methods described herein, wherein the experimental DNA comprises (a) a promoter from the transcriptional regulator; and (b) promoters from the plurality of transcriptional regulators; wherein a transcriptional regulatory network is identified if the transcriptional regulator regulates itself or if it regulates at least one of the plurality of transcriptional regulators.

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The invention further provides a method of identifying transcriptional regulatory networks in a cell, the method comprising (a) determining, by repeating a

method of identifying the targets of transcriptional regulator for each of a plurality of transcriptional regulators, the genes in a subset which are regulated by each of the plurality of transcriptional regulators, wherein the experimental DNA comprises promoter regions for each of the plurality of transcriptional regulators; (b) determining if any one of the plurality of transcriptional regulators are regulated by at least one of the plurality of transcriptional regulators; wherein a transcriptional regulatory network is identified if any one of the plurality of transcriptional regulators is regulated by at least one of the plurality of transcriptional regulators.

The invention also provides a DNA microarray for determining promoter occupancy in a human cell, the microarray comprising (1) at least 10,000 experimental spots, each experimental spot comprising an experimental DNA, each experimental DNA comprising a promoter region from a human gene in the subset; and (2) at least 100 control spots, each control spot comprising a control DNA, each control DNA comprising a non-promoter region; wherein at least 75% of the promoter regions comprise from at least 700bp upstream to at least 200 bp downstream of the transcriptional start site.

Another aspect of the invention provides a method of estimating if a transcriptional regulator is a global transcriptional regulator, the method comprising (a) selectively isolating chromatin from a tissue; (b) identifying promoter regions from the chromatin which are bound by a candidate global transcriptional regulator; (c) identifying promoter regions from the chromatin which are bound by a member of the basal transcriptional machinery; and (d) comparing the promoter regions identified in steps (b) and (c) to determine the ratio between (i) the number of promoter regions bound by both the candidate global transcriptional regulator and the member of the basal transcriptional machinery; and (ii) the number of promoter regions bound by the member of the basal transcriptional machinery, wherein a transcriptional regulator is a global transcriptional regulator when the ratio is greater than 0.2.

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The invention further provides methods of identifying targets for therapeutics.

In one aspect, the invention provides a method of identifying at least one target gene for

the development of a therapeutic to treat or prevent a disorder in a subject, wherein at least one form of the disorder is caused by an altered activity in a transcriptional regulator or in a suspected transcriptional regulator, the method comprising (a) identifying the genes regulated by the transcriptional regulator in a cell; (b) determining if the transcriptional regulator is a broad-acting transcriptional regulator or a narrow-acting transcriptional regulator, wherein if the transcriptional regulator is a broad acting transcriptional regulator then the transcriptional regulator is a target gene for the development of a therapeutic, and wherein if the transcriptional regulator is a narrow acting transcriptional regulator then (i) determining if at least one gene regulated by the transcriptional regulator is likely causative in the disorder, wherein a gene that is likely causative in the disorder is a target gene for the development of a therapeutic; and (ii) reiterating steps (a) and (b) for at least one gene that is regulated by the transcriptional regulator in the cell and that either (1) encodes a transcriptional regulator or (2) is suspected to encode a transcriptional regulator, with the modification that the transcriptional regulator of steps (a) and (b) is said gene, thereby identifying at least one target gene for the development of a therapeutic to treat or prevent a disorder in the subject.

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The invention also provides methods of treating or preventing disease. In one aspect, the invention provides a method of treating or preventing type II diabetes in a subject, comprising administering to the subject a therapeutically effective amount of an agent that increases the global transcriptional activity of HNF4alpha.

In another aspect, the invention provides a method of treating or preventing a disorder associated with low transcriptional activity of HNF4alpha in a subject, comprising administering to the subject a therapeutically effective amount of an agent that increases the global transcriptional activity of HNF4alpha. A related aspect provides a method of treating or preventing a disorder associated with high transcriptional activity of HNF4alpha in a subject, comprising administering to the subject a therapeutically effective amount of an agent that decreases the global transcriptional activity of HNF4alpha.

The invention also provides a method of increasing the global transcriptional activity in a liver or a pancreatic cell comprising contacting the cell with an agent which increases the global transcriptional activity of HNF4alpha. A related aspect provides a method of decreasing the global transcriptional activity in a liver or a pancreatic cell comprising contacting the cell with an agent which decreases the global transcriptional activity of HNF4alpha.

One aspect of the invention provides methods of regulating the expression level of genes. On aspect provides a method of regulating the expression level of any one of the genes in Figure 13 in a hepatocyte, the method comprising contacting the cell with an agent which regulates the transcriptional activity of HNF1alpha. A related aspect provides a method of regulating the expression level of any one of the genes in Figure 14 in a pancreatic cell, the method comprising contacting the cell with an agent which regulates the transcriptional activity of HNF1alpha.

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Another aspect of the invention provides a method of regulating the expression level of any one of the genes in Figure 16 in a hepatocyte, the method comprising contacting the cell with an agent which regulates the transcriptional activity of HNF6. A related aspect provides a method of regulating the expression level of any one of the genes in Figure 17 in a pancreatic cell, the method comprising contacting the cell with an agent which regulates the transcriptional activity of HNF6.

Yet another aspect of the invention provides a method of regulating the expression level of any one of the genes in Figure 18 in a hepatocyte, the method comprising contacting the cell with an agent which regulates the transcriptional activity of HNF4alpha. A related aspect provides a method of regulating the expression level of any one of the genes in Figure 19 in a pancreatic cell, the method comprising contacting the cell with an agent which regulated the transcriptional activity of HNF4alpha.

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The invention also provides methods for identifying transcriptionally active genes that are regulated by a transcriptional regulator in a cell. In one aspect, the

invention provides a method of identifying transcriptionally active genes that are regulated by a transcriptional regulator in a cell, the method comprising (a) selectively isolating chromatin from a tissue; (b) identifying promoter regions from the chromatin that are bound by the transcriptional regulator; (c) identifying promoter regions from the chromatin that are bound by a member of the basal transcriptional machinery; and (d) comparing the promoter regions identified in steps (b) and (c) to determine overlapping genes, wherein the overlapping genes are transcriptionally active genes regulated by the transcriptional regulator.

10 BRIEF DESCRIPTION OF THE DRAWINGS

Figures 1A-1C show genome-scale location analysis of HNF regulators in human tissues. (A) Hepatocytes and pancreatic islets were obtained from tissue distribution programs. These cells were treated with formaldehyde to covalently link transcription factors to DNA sites of interaction. Cells were harvested, and chromatin in cell lysates was sheared by sonication. The regulator-DNA complexes were enriched by chromatin immunoprecipitation with specific antibodies, the crosslinks were reversed, and enriched DNA fragments and control genomic DNA fragments were amplified using ligation-mediated PCR. The amplified DNA preparations, labeled with distinct fluorophores, were mixed and hybridized onto a promoter array. (B) Venn diagram showing the overlap of HNF1α, HNF6, and HNF4α bound promoters in hepatocytes (top) and pancreatic islets (bottom). (C) The collection of genes occupied by RNA polymerase II in hepatocytes is displayed as a circle, with the genes bound by HNF1α, HNF6, and HNF4α outlined collectively as a fraction of the chart. The relative contributions of HNF1α, HNF6, and HNF4α are shown as framing arcs.

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Figures 2A-2B show transcriptional regulatory networks and motifs. (A) HNF1α, HNF6, and HNF4α are at the center of tissue-specific transcriptional regulatory networks. In these examples selected for illustration, regulatory proteins and their gene targets are represented as circles and boxes, respectively. Solid arrows indicate protein-DNA interactions, and genes encoding regulators are linked to their protein products by dashed lines. The HNF4a7 promoter, also known as the P2 promoter (24, 25), was recently implicated as a major human diabetes susceptibility locus (see text). (B)

Examples of regulatory network motifs in hepatocytes. For instance, in the multi-component loop, HNF1 α protein binds to the promoter of the HNF4 α gene, and the HNF4 α protein binds to the promoter of the HNF1 α gene. These network motifs were uncovered by searching binding data with various algorithms; for details on the algorithms used and a full list of motifs found, see (20).

Figure 3 shows one embodiment of a strategy for the identification of at least one target gene of a master regulator for the development of a therapeutic to treat or prevent a disorder.

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Figure 4 shows a Venn diagram showing the overlap of two single, independent ChIP experiments using hepatocytes with anti-HNF4a antibodies sc-6556 and sc-8987.

Figure 5 shows a Western blot of HNF4a in HepG2 cells using 50 µg of cell lysate protein with Ab sc-6556. The lower running band is approximately 50 kDa, which is the canonical molecular weight for HNF4a, and the higher running band is the appropriate location for HNF4a dimer. A very similar gel showing HNF4a antibody specificity for sc-6556 is available at the Santa Cruz website (www.scbt.com).

Figures 6A-6D show scatterplots of attempted chromatin immunoprecipitations performed with the anti-HNF4a antibody sc-6556 using Jurkat (T-lymphocyte derived, 6A), BJ-T (foreskin fibroblast derived, 6B), and U937 (histocyte derived, 6C) cells. To demonstrate the noise inherent in the array analysis, applicants show a scatterplot of a sample of input DNA, split, labeled with the two fluorophores, and hybridized to an array (6D). Identical control experiments performed using the anti-HNF1a antibody sc-6547 afforded essentially identical results.

Figure 7 shows a scatterplot of a chromatin immunoprecipitation performed with preimmune commercial rabbit serum using hepatocytes (left). Goat pre-immune serum and two rabbit sera from different individuals gave a similar scatterplot. For comparison, applicants show the scatterplot for an equivalent ChIP with the anti-HNF4a antibody sc-6556 using hepatocytes (right).

Figure 8 shows a Venn diagram showing the overlap of the sets of promoters bound by HNF4α and RNA Pol II in hepatocytes and pancreatic islets.

Figure 9 shows a composite gel of gene-specific chromatin immunoprecipitation reactions using anti-HNF4α antibody sc-6556 with crosslinked human hepatocytes.

Figure 10 shows composite gel of gene-specific chromatin immunoprecipitation reactions using anti-HNF1α antibody sc-6547 with crosslinked human hepatocytes.

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Figure 11 shows a partial list of proximal promoters occupied by of HNF1a in human hepatocytes and pancreatic islets. These genes were assigned to functional categories using the program ProtoGo; genes not in this automated GO ontology database were assigned using Locuslink information. Four genes are shown for each tissue/category combination; for some combinations, fewer than 4 promoters qualified as targets. Hypothetical and functionally uncharacterized genes are not shown. A complete list of targets is available in Figures 13 and 14.

Figure 12 shows Occupancy of BJ-T and tissue-specific promoter sets by HNF factors.

(*) Indicates that comparisons between BJ-T and primary tissues used only a subset of Hu13K array promoters, as RNA Pol II was profiled in BJ-T cells using a smaller, prototype array. The denominator in the above fractions represents the number of targets the HNF factor of interest occupied in the set of RNA Pol II occupied promoters that are either BJ-T specific or primary tissue specific.

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Figure 13 shows HNF1\alpha bound promoters in hepatocytes

Figure 14 shows HNF1a bound promoters in pancreatic islets.

Figures 15A-15D show genes previously suggested to be regulated by HNF1a and HNF4a. 'Direct' binding is in vivo ChIP and in vivo footprinting, 'in vitro' binding is primarily gel mobility retardation assays and in vitro footprinting, and 'indirect' is

primarily transient transfections. 'Sequence-based' uses a number of different criteria to qualify binding. Note that some duplicate reports are omitted, as are a handful of recent large-scale screens, (e.g. Tronche 1997, Shih 2001, etc.).

5 Figure 16 shows HNF6 bound promoters in hepatocytes.

Figure 17 shows HNF6 bound promoters in pancreatic islets.

Figure 18A-18C show HNF4α bound promoters in hepatocytes.

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Figures 19A-19C show HNF4a bound promoters in pancreatic islets.

Figures 20A-20B show the feed forward regulatory motifs in hepatocytes. The regulatory modules here were derived as described in exemplification. Feed forwards only involving HNF1a and HNF4a are also multi-input motifs, as they bind each other's promoters in a multicomponent loop.

Figures 21A-21B show multi-input motifs in hepatocytes. The regulatory modules here were derived as described in the exemplification. MIMs for the HNF6/HNF4a and HNF1a/HNF4a are listed in Figure 20 as feedforward motifs.

Figures 22A-22B show the feed forward regulatory motifs in pancreatic islets. The regulatory modules here were derived as described in Supporting Online Material. Feed forwards only involving HNF1a and HNF4a are also multiinput motifs, as they bind each other's promoters in a multicomponent loop.

Figures 23A-23B show multi-Input motifs in pancreatic islets. The regulatory modules here were derived as described in Supporting Online Material. MIMs for the HNF6/HNF4a and HNF1a/HNF4a are listed in Figure 22 as feedforward.

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Figures 24A-24B show transcriptional regulators occupied by HNF1a and HNF4a. Network of DNA regulators downstream of HNF1a and HNF4a in hepatocytes and

islets. Target genes that are among the Gene Ontology "DNA-regulators" category were compiled, and are listed according to functional subcategory.

DETAILED DESCRIPTION OF THE INVENTION

5 I. Overview

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In certain aspects, the invention provides methods related to transcriptional regulators. Some aspects of the invention provide methods for the identification of genes whose transcription is regulated by a specific transcriptional regulator in a cell. Some of these methods comprise determining the promoter occupancy of the transcriptional regulator using a combination of chromatin immunoprecipitation and/or DNA microarray analysis of the promoter regions that are physically associated with the transcriptional regulator in the cell. In some embodiments of the methods described herein, the DNA microarray comprises both experimental spots containing promoter DNA, and control spots containing non-promoter DNA. The methods described herein may be applied to any cell type, including transplant grade primary human tissue. Furthermore, the method described herein can be used to compare the function of transcriptional regulators across cell types, or across two populations, such as healthy and disease-afflicted subjects.

In a related aspect, the invention provides methods of identifying regulatory networks, or pathways. Some methods comprise identifying the transcriptional regulators which are regulated by a given transcriptional regulator, and optionally, determining the genes that are regulated by those transcriptional regulators. Pathways that may be identified using the methods described herein include autoregulatory, multicomponent, feed-forward, and multi-components loops, as well as regulatory chains.

The invention also provides methods of determining if a transcriptional regulator is a global transcriptional regulator. In some aspects, such methods comprise determining the promoter occupancy of both a transcriptional regulator and a member of the basal transcriptional machinery. Comparison of the promoter occupancy by the transcriptional regulator and by the member of the basal transcriptional machinery

allows the identification of transcriptionally active promoters that are bound and regulated by the transcription regulator. Other methods further comprise extrapolating from the set of promoters that were examined to the total number of promoters in the genome to determine the approximate number of transcriptionally active promoters in a cell that are under the control of a specific transcriptional factor or to determine if the transcriptional regulator is a global transcriptional regulator.

Other aspects of the invention provide methods of identifying therapeutic targets to treat disease. One specific aspect of the invention relates to identifying at least one target gene for the development of a therapeutic agent to treat or prevent a disorder in a subject, preferably a disorder in which at least one form of the disorder is caused by an altered activity in a transcriptional regulator or in a gene suspected to encode a transcriptional regulator. Some of the methods provided herein to identify therapeutic targets comprise determining if a transcriptional regulator implicated in the disease is a broad-acting or a narrow-acting transcriptional regulator, such as by identifying at least a subset of the genes that it regulates in a cell, wherein broad-acting transcriptional regulators are targets for therapeutic agents. If the transcriptional regulator is narrow-acting, then the genes that it regulates may be examined further to determine if any are broad-acting transcriptional regulators (for those genes encoding transcriptional regulators) or if any of the genes are causative to the disease state *i.e.* they regulate a pathway or network that is impaired in the disease state.

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The invention further provides methods for the treatment of disease. Some aspects of the invention provide methods of treating metabolic disorders, such as type II diabetes. Specific aspects of the invention provide methods of treating or preventing type II diabetes in a subject by administering to the subject a therapeutically effective amount of an agent that increases the global transcriptional activity of HNF4α. Furthermore, the invention provides methods for modulating the expression level of genes. Such methods are based, in part, on the finding by Applicants of genes which are transcriptionally regulated by HNF1α, HNF4α or HNF6 in hepatocytes and pancreatic cells. In a related aspect, the invention provides methods of modulating and expression level of, and alleviating a disease state associated with the abnormal

expression of, the genes in Figures 13-19 by modulating the transcriptional activity or expression of HNF1 α , HNF4 α or HNF6. In specific embodiments, the expression of the genes is modulated in hepatocytes, pancreatic cells, or both.

5 II. Definitions

For convenience, certain terms employed in the specification, examples, and appended claims, are collected here. Unless defined otherwise, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention belongs.

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The articles "a" and "an" are used herein to refer to one or to more than one.

(i.e., to at least one) of the grammatical object of the article. By way of example, "an element" means one element or more than one element.

The term "including" is used herein to mean, and is used interchangeably with, the phrase "including but not limited" to.

The term "or" is used herein to mean, and is used interchangeably with, the term "and/or," unless context clearly indicates otherwise.

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The term "such as" is used herein to mean, and is used interchangeably, with the phrase "such as but not limited to".

A "patient" or "subject" to be treated by the method of the invention can mean either a human or non-human animal, preferably a mammal.

The terms "alpha" and " α " are used interchangeably, as are the terms "beta" and " β ".

The term "encoding" comprises an RNA product resulting from transcription of a DNA molecule, a protein resulting from the translation of an RNA molecule, or a protein resulting from the transcription of a DNA molecule and the subsequent

translation of the RNA product.

A "promoter" is a nucleic acid sequence that directs transcription of a nucleic acid. A promoter includes nucleic acid sequences near the start site of transcription, e.g., a TATA box, see, e.g., Butler and Kadonaga (2002) Genes Dev. 16:2583-2592; Georgel (2002) Biochem. Cell Biol. 80:295-300. A promoter also optionally includes distal enhancer or repressor elements, which can be located as much as several thousand base pairs on either side from the start site of transcription. A "constitutive" promoter is a promoter that is active under most environmental and developmental conditions, while an "inducible", promoter is a promoter is active or activated under, e.g., specific environmental or developmental conditions.

The term "expression" is used herein to mean the process by which a polypeptide is produced from DNA. The process involves the transcription of the gene into mRNA and the translation of this mRNA into a polypeptide. Depending on the context in which used, "expression" may refer to the production of RNA, protein or both.

The term "recombinant" is used herein to mean any nucleic acid comprising sequences which are not adjacent in nature. A recombinant nucleic acid may be generated *in vitro*, for example by using the methods of molecular biology, or *in vivo*, for example by insertion of a nucleic acid at a novel chromosomal location by homologous or non-homologous recombination.

The term "transcriptional regulator" refers to a biochemical element that acts to prevent or inhibit the transcription of a promoter-driven DNA sequence under certain environmental conditions (e.g., a repressor or nuclear inhibitory protein), or to permit or stimulate the transcription of the promoter-driven DNA sequence under certain environmental conditions (e.g., an inducer or an enhancer).

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The term "microarray" refers to an array of distinct polynucleotides or oligonucleotides synthesized on a substrate, such as paper, nylon or other type of

membrane, filter, chip, glass slide, or any other suitable solid support.

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The terms "disorders" and "diseases" are used inclusively and refer to any deviation from the normal structure or function of any part, organ or system of the body (or any combination thereof). A specific disease is manifested by characteristic symptoms and signs, including biological, chemical and physical changes, and is often associated with a variety of other factors including, but not limited to, demographic, environmental, employment, genetic and medically historical factors. Certain characteristic signs, symptoms, and related factors can be quantitated through a variety of methods to yield important diagnostic information.

The terms "level of expression of a gene in a cell" or "gene expression level" refer to the level of mRNA, as well as pre-mRNA nascent transcript(s), transcript processing intermediates, mature mRNA(s) and degradation products, encoded by the gene in the cell.

The term "modulation" refers to upregulation (i.e., activation or stimulation), downregulation (i.e., inhibition or suppression) of a response, or the two in combination or apart. A "modulator" is a compound or molecule that modulates, and may be, e.g., an agonist, antagonist, activator, stimulator, suppressor, or inhibitor.

The term "agonist" refers to an agent that mimics or up-regulates (e.g., potentiates or supplements) the bioactivity of a protein, e.g., polypeptide X. An agonist may be a wild-type protein or derivative thereof having at least one bioactivity of the wild-type protein. An agonist may also be a compound that upregulates expression of a gene or which increases at least one bioactivity of a protein. An agonist may also be a compound which increases the interaction of a polypeptide with another molecule, e.g., a target peptide or nucleic acid.

The term "antagonist" refers to an agent that downregulates (e.g., suppresses or inhibits) at least one bioactivity of a protein. An antagonist may be a compound which inhibits or decreases the interaction between a protein and another molecule, e.g., a

target peptide or enzyme substrate. An antagonist may also be a compound that downregulates expression of a gene or which reduces the amount of expressed protein present.

The term "prophylactic" or "therapeutic" treatment refers to administration to the subject of one or more of the subject compositions. If it is administered prior to clinical manifestation of the unwanted condition (e.g., disease or other unwanted state of the host animal) then the treatment is prophylactic, i.e., it protects the host against developing the unwanted condition, whereas if administered after manifestation of the unwanted condition, the treatment is therapeutic (i.e., it is intended to diminish, ameliorate or maintain the existing unwanted condition or side effects therefrom).

The term "therapeutic effect" refers to a local or systemic effect in animals, particularly mammals, and more particularly humans caused by a pharmacologically active substance. The term thus means any substance intended for use in the diagnosis, cure, mitigation, treatment or prevention of disease or in the enhancement of desirable physical or mental development and conditions in an animal or human. The phrase "therapeutically-effective amount" means that amount of such a substance that produces some desired local or systemic effect at a reasonable benefit/risk ratio applicable to any treatment. In certain embodiments, a therapeutically-effective amount of a compound will depend on its therapeutic index, solubility, and the like. For example, certain compounds discovered by the methods of the present invention may be administered in a sufficient amount to produce a reasonable benefit/risk ratio applicable to such treatment.

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A probe that is "labeled" is detectable, either directly or indirectly, by spectroscopic, photochemical, biochemical, immunochemical, isotopic, or chemical means. For example, useful labels include ³²P, ³³P, ³⁵S, ¹⁴C, ³H, ¹²⁵I, stable isotopes, fluorescent dyes and fluorettes (Rozinov and Nolan (1998) Chem. Biol 5:713-728; Molecular Probes, Inc. (2003) Catalogue, Molecular Probes, Eugene Oreg.), electrondense reagents, enzymes and/or substrates, e.g., as used in enzyme-linked immunoassays as with those using alkaline phosphatase or horse radish peroxidase. The

label or detectable moiety is typically bound, either covalently, through a linker or chemical bound, or through ionic, van der Waals or hydrogen bonds to the molecule to be detected. "Radiolabeled" refers to a compound to which a radioisotope has been attached through covalent or non-covalent means. A "fluorophore" is a compound or moiety that absorbs radiant energy of one wavelength and emits radiant energy of a second, longer wavelength.

A "labeled nucleic acid probe or oligonucleotide" is one that is bound, either covalently, through a linker or a chemical bond, or noncovalently, through ionic, van der Waals, electrostatic, or hydrogen bonds to a label such that the presence of the probe can be detected by detecting the presence of the label bound to the probe. The probes are preferably directly labeled as with isotopes, chromophores, fluorophores, chromogens, or indirectly labeled such as with biotin to which a streptavidin complex or avidin complex can later bind.

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A "nucleic acid probe" is a nucleic acid capable of binding to a target nucleic acid of complementary sequence, usually through complementary base pairing, e.g., through hydrogen bond formation. A probe may include natural, e.g., A, G, C, or T, or modified bases, e.g., 7-deazaguanosine, inosine, etc. The bases in a probe can be joined by a linkage other than a phosphodiester bond. Probes can be peptide nucleic acids in which the constituent bases are joined by peptide bonds rather than phosphodiester linkages. It will be understood by one of skill in the art that probes may bind target sequences lacking complete complementarity with the probe sequence depending upon the stringency of the hybridization conditions.

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"Small molecule" is defined as a molecule with a molecular weight that is less than 10 kD, typically less than 2 kD, and preferably less than 1 KD. Small molecules include, but are not limited to, inorganic molecules, organic molecules, organic molecules containing an inorganic component, molecules comprising a radioactive atom, synthetic molecules, peptide mimetics; and antibody mimetics. As a therapeutic, a small molecule may be more permeable to cells, less susceptible to degradation, and less apt to elicit an immune response than large molecules. Small molecule toxins are

described, see, e.g., U.S. Pat. No. 6,326,482 issued to Stewart, et al.

A small molecule refers to a composition, which has a molecular weight of less than about 1000 kDa.

5 III. Identification of Transcriptional Targets and Transcriptional Networks

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One aspect of the invention provides a method of determining which genes from a subset of genes are regulated by a transcriptional regulator in a cell, the method comprising (a) selectively isolating chromatin from a cell which expresses the transcriptional regulator to generate isolated chromatin; (b) selectively isolating chromatin fragments from the isolated chromatin to generate bound chromatin fragments, wherein the bound chromatin fragments are bound by the transcriptional regulator; (c) amplifying both the bound chromatin fragments to generate amplified chromatin fragments and the isolated chromatin to generate amplified control chromatin; (d) hybridizing the amplified control chromatin and the amplified chromatin fragments to a DNA microarray, wherein the DNA microarray comprises (1) at least 10,000 experimental spots, each experimental spot comprising an experimental DNA, each experimental DNA comprising a promoter region from a gene in the subset; and (2) at least 100 control spots, each control spot comprising a control DNA, each control DNA comprising a non-promoter region; and (e) determining and comparing a hybridization signal at each of the spots on the microarray between those generated by (1) the amplified control chromatin; and (2) the amplified chromatin fragments; wherein a gene in the subset is said to be regulated by the transcriptional regulator in the cell if a spot comprising a promoter region of said gene displays a higher level of hybridization by the amplified chromatin fragments than by the amplified control chromatin.

Methods of isolating chromatin, and in particular chromatin fragments that are bound by a transcriptional regulator, may be carried out by any method known to one skilled in the art, including by cross-linking the transcriptional regulator to chromatin, fragmenting the chromatin, and immunoprecipitating the transcriptional regulators.

In a preferred embodiment, the chromatin fragments bound by the

transcriptional regulator are isolated using chromatin immunoprecipitation (ChIP). Briefly, this technique involves the use of a specific antibody to immunoprecipitate chromatin complexes comprising the corresponding antigen *i.e.* the transcriptional regulator, and examination of the nucleotide sequences present in the immunoprecipitate. Immunoprecipitation of a particular sequence by the antibody is indicative of interaction of the antigen with that sequence. See, for example, O'Neill et al. in *Methods in Enzymology*, Vol. 274, Academic Press, San Diego, 1999, pp. 189-197; Kuo et al. (1999) *Method* 19:425-433; and Ausubel et al., supra, Chapter 21.

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In one embodiment, the chromatin immunoprecipitation technique is applied as follows. Cells which express the transcriptional regulator of interest, such as a native transcriptional regulator or a recombinant transcriptional regulator, are treated with an agent that crosslinks the transcriptional regulator to chromatin if that transcriptional regulator is stably bound to it. In one embodiment of the methods described herein, the crosslinking is formaldehyde crosslinking (Solomon, M.J. and Varshavsky, A., Proc. Natl. Sci. USA 82:6470-6474; Orlando, V., TIBS, 25:99-104). UV light may also be used (Pashev et al. *Trends Biochem Sci.* 1991;16(9):323-6; Zhang L et al. *Biochem Biophys Res Commun.* 2004;322(3):705-11).

Subsequent to crosslinking, cellular nucleic acid is isolated, sheared such as by sonication and incubated in the presence of an antibody directed against the transcriptional regulator. Antibody-antigen complexes are precipitated, crosslinks are reversed (for example, formaldehyde-induced DNA-protein crosslinks can be reversed by heating) so that the sequence content of the immunoprecipitated DNA is tested for the presence of a specific sequence, for example, promoter regions. The antibody may bind directly to an epitope on the transcriptional regulator or it may bind to a tag on the regulator, such as a myc tag when used with an anti-Myc antibody (Santa Cruz Biotechnology, sc-764).

In yet another embodiment, a non-antibody agent with affinity for the transcriptional regulator or for a tag used to it is used in place of the antibody. For example, if the transcriptional regulator comprises an affinity tag, such as a six-

histidine tag, complexes may be isolated by affinity chromatography to nickel-containing sepharose. Additional variations on ChIP methods within the scope of the invention may be found in Kurdistani et al. Methods. 2003 31(1):90-5; O'Neill et al. Methods. 2003, 31(1):76-82; Spencer et al., Methods. 2003;31(1):67-75; and Orlando et al. Methods 11: 205-214 (1997).

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In an alternate embodiment of the methods described herein for identifying genes regulated by a transcriptional regulator, amplified chromatin fragments from a control immunoprecipitation reaction are used in place of the isolated chromatin as a control. For example, an antibody that does not react with the transcription factor being tested may be used in a chromatin IP procedure to isolate control chromatin, which can then be compared to the chromatin isolated using an antibody that does react with the transcriptional regulator. In preferred embodiments, the antibody that does not react with the transcription factor being tested also does not react with other transcriptional regulators or DNA binding proteins.

In one embodiment, the amplified control chromatin and the amplified chromatin fragments are generated from their corresponding template DNA using ligation-mediated polymerase chain reaction (LM-PCR) (e.g., see Current Protocols in Molecular Biology, Ausubel, F. M. et al., eds. 1991, and U.S. Application No. 2003/0143599, the teachings of which are incorporated herein by reference) in their entirety. In specific embodiments, LM-PCR comprises fluorescently labeling amplified DNA by including fluorescently-tagged nucleotides in the LM-PCR reaction. Additional variations for manipulating and examining chromatin using microarrays have described in U.S. Patent Nos. 6,410,243, the teachings of which are incorporated herein by reference.

In one embodiment, the labelled or unlabeled probes are hybridized to DNA microarray, such as is described in U.S. Patent No. 6,410,243. Microarrays, also called "biochips" or "arrays" are miniaturized devices typically with dimensions in the micrometer to millimeter range for performing chemical and biochemical reactions and are particularly suited for embodiments of the invention. Arrays may be constructed via

microelectronic and/or microfabrication using essentially any and all techniques known and available in the semiconductor industry and/or in the biochemistry industry, provided only that such techniques are amenable to and compatible with the deposition and screening of polynucleotide sequences. Microarrays are particularly desirable for their virtues of high sample throughput and low cost for generating profiles and other data. Additional variations for manipulating and examining chromatin using microarrays have described in U.S. Patent Nos. 6,410,243, the teachings of which are incorporated herein by reference.

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In one embodiment of the methods described, amplified control chromatin and the amplified chromatin fragments are hybridized to a DNA microarray that includes experimental spots that represent all or a subset (e.g., a chromosome or chromosomes) of the genome. The fluorescent intensity of each experimental spot on the microarray from the amplified chromatin fragments relative to the amplified control chromatin indicates whether the protein of interest is bound to the DNA region located at that particular spot. Hence, the methods described herein allow the detection of protein-DNA interactions across an entire genome.

In some embodiments of the methods described herein, the promoter region of a gene comprises from at least 700bp upstream to at least 200 bp downstream of the transcriptional start site of the gene. In some embodiments, the promoter region comprises at least about 30, 40, 50, or 60 nucleotides in length. In specific embodiments, the promoter region of a gene as found on the spots of the microarray comprises a sequence of at least 30 nucleotides whose sequence is identical to a region stretching from 3 kb upstream to 1 kb downstream of the transcriptional start site of said gene. In some embodiments, the DNA microarray includes control spots of non-promoter DNA. In specific embodiment, the non-promoter region comprises an open reading frame. In preferred embodiments, the non-promoter regions comprise genomic regions which are not bound by transcriptional regulators, and preferably which are not bound by the transcriptional regulator being tested. In some embodiments, not all the experimental spots or the control spots comprise experimental DNA or control DNA, respectively. Furthermore, in some specific embodiments some spots comprise control

DNA which comprises promoter DNA. One skilled in the art may determine the number of experimental or control spots for a given application.

In some embodiments of the methods described herein, the level of hybridization of the amplified chromatin fragments to each experimental spot is normalized by the level of hybridization of the amplified chromatin fragments to the control spots. In specific embodiments, the normalization is performed by subtracting the mean level of hybridization of the amplified chromatin fragments to the control spots from the level of hybridization of the amplified chromatin fragments at each experimental spot.

Methods of analyzing data from microarrays are well-described in the art, including in DNA Microarrays: A Molecular Cloning Manual, Ed by Bowtel and Sambrook (Cold Spring Harbor Laboratory Press, 2002); Microarrays for an Integrative Genomics by Kohana (MIT Press, 2002); A Biologist's Guide to Analysis of DNA Microarray Data, by Knudsen (Wiley, John & Sons, Incorporated, 2002); and DNA Microarrays: A Practical Approach, Vol. 205 by Schema (Oxford University Press, 1999); and Methods of Microarray Data Analysis II, ed by Lin et al. (Kluwer Academic Publishers, 2002), hereby incorporated by reference in their entirety.

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In some embodiments of any of the methods described herein, the transcriptional regulator is native to the cell. By native it is meant that the transcriptional regulator naturally occurs in the cell. In other embodiments, the transcriptional regulator is a recombinant transcriptional regulator. In some embodiments, the transcriptional regulator originates from a species which is different from that of the cell. In some embodiments, the transcriptional regulator is a viral transcriptional regulator. In such embodiments, a cell may be contacted with a virus and chromatin extracted from the infected cell after allowing sufficient time for the viral proteins to be expressed. In some embodiments, recombinant transcriptional regulators have missense mutations, truncations, or inserted sequences or entire domains from other naturally occurring proteins. A tagged recombinant transcriptional regulator may be used in some embodiments the methods of the present invention as

the tag may facilitate the immunoprecipitation of the regulator.

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In certain embodiments of the invention, transcriptional regulators comprise specific transcription factors, coactivators, corepressors or complexes thereof. Transcription factors bind to specific cognate DNA elements such as promoters, enhancers and silencer elements, and are responsible for regulating gene expression. Transcription factors may be activators of transcription, repressors of transcription or both, depending on the cellular context. Transcription factors may belong to any class or type of known or identified transcription factor. Examples of known families or structurally-related transcription factors include helix-loop-helix, leucine zipper, zinc finger, ring finger, and hormone receptors. Transcription factors may also be selected based upon their known association with a disease or the regulation of one or more genes. For example, transcription factors such as c-myc, Rel/Nf-kB, neuroD, c-fos, cjun, and E2F may be targeted. Antibodies directed to any transcriptional coactivator or corepressor may also be used according to the invention. Examples of specific coactivators include CBP, CTIIA, and SRA, while specific examples of corepressors include the mSin3 proteins, MITR, and LEUNIG. Furthermore, the genes regulated by proteins associated with transcriptional complexes, such as the histone acetylases (HATs) and histone deacetylases (HDACs), may also de determined using the methods described herein.

In one embodiment of the methods described herein, the cell is a primary cell. Primary cells are directly isolated from an organism and have undergone minimum passaging *in vitro*, and thus maintain most of the phenotypic characteristics of cells in the organism. In a specific embodiment, the primary cells are primary cells that have doubled less than 10 times *ex vivo*. In some embodiments, the cell is derived from transplant grade tissue or freshly isolated tissue. The cell type used in the assays described herein may be any cell type. The cell may be eukaryotic or prokaryotic, from a metazoan or from a single-celled organism such as yeast. In some preferred embodiments the cell is a mammalian cell, such as a cell from a rodent, a primate or a human. The cell may be a wild-type cell or a cell that has been genetically modified by recombinant means or by exposure to mutagens. The cell may be a transformed cell or

an immortalized cell. In some embodiments, the cell is from an organism afflicted by a disease. In some embodiments, the cell comprises a genetic mutation that results in disease, such as in a hyperplastic condition.

In some embodiments, the cell is derived from transplant-grade tissue or freshly isolated tissue. In some embodiments, the cell is derived from a tissue biopsy, such as from a subject afflicted with, or suspected of being afflicted with, a disorder. In another embodiment, the cell is isolated from a bodily fluid or bodily secretion, including serum, plasma, saliva, tears, sweat, semen, amniotic fluid, vaginal secretions, nasal secretions, synovial fluid, spinal fluid, phlegm, bronchoalveolar lavage fluid, blister fluid, pus, stool and intracranial fluid. The cell may be a live cell or a cell that has been preserved, such as by treatment with formalin, B5, Zenker's fixatives, Lugol's solution, Carnoy's Fixative, F13 fixative, or other preservatives, or a cell that has been preserved by freezing.

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In some embodiments of the methods described herein, the cell has been treated with an agent, such as compound or a drug, prior to isolation of chromatin. Some preferred agents include those which bind to or regulate the expression of transcriptional regulators. In some embodiments, the genes that are regulated by a given transcriptional regulator are determined both in a cell that is contacted with an agent and in a cell that is not contacted with the agent, or that is contacted with a different amount of the agent. Such methods may be used to identify compounds that alter the types of genes and/or the extent to which a transcriptional regulators controls transcription of those genes. Furthermore, such approaches may be used to screen for agents which alter the activity, specificity or expression of a transcriptional regulator.

In some embodiment of the methods described herein for identifying genes regulated by a transcriptional regulator, a higher level of hybridization by the amplified chromatin fragments than by the amplified control chromatin comprises at least a two-fold higher level of hybridization. The threshold for what constitutes a higher level of hybridization, may be adjusted by one skilled in the art for the particular application. Higher levels of hybridization are expected to yield a smaller target size but with higher

certainty that a given gene above that threshold is regulated by the transcriptional regulator in that cell in vivo.

In other embodiments of the methods described herein for identifying genes regulated by a transcriptional regulator, the transcriptional regulator is a basal transcription factor or a component of the basal transcription machinery. In specific embodiments, components of the basal transcription machinery comprise RNA polymerases, including poll, poll and pollil, TBP, NTF-1 and Sp1 and any other component of TFIID, including, for example, the TAFs (e.g. TAF250, TAF150, TAF135, TAF95, TAF80, TAF55, TAF31, TAF28, and TAF20), or any other component of a polymerase holoenzyme.

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Another aspect of the invention provides a method of identifying transcriptionally active genes that are regulated by a transcriptional regulator in a cell. The method comprises determining what genes are regulated by the transcriptional regulator and determining which ones are transcriptionally active in the cell. In one embodiment, a set of genes which are transcriptionally active is the set of genes whose promoters are bound by an RNA polymerase, such as RNA polymerase II, or by a member of the basal transcription machinery. Alternatively, genes which are transcriptionally active may be identified using other techniques know in the art. For example, mRNA from a cell which expresses the transcriptional regulator can be collected and examined on a DNA microarray which comprises coding sequences in order to determine which genes are being transcribed.

In one embodiment, the invention provides a method of identifying transcriptionally active genes that are regulated by a transcriptional regulator in a cell, the method comprising (a) selectively isolating chromatin from a tissue; (b) identifying promoter regions from the chromatin that are bound by the transcriptional regulator; (c) identifying promoter regions from the chromatin that are bound by a member of the basal transcriptional machinery; and (d) comparing the promoter regions identified in steps (b) and (c) to determine overlapping genes, wherein the overlapping genes are transcriptionally active genes regulated by the transcriptional regulator.

In a related aspect, the invention provides methods to determine if a transcriptional regulator is a global transcription regulator. One method comprises estimating if a transcriptional regulator is a global transcriptional regulator, the method comprising (a) selectively isolating chromatin from a tissue; (b) identifying promoter regions from the chromatin which are bound by a candidate global transcriptional regulator; (c) identifying promoter regions from the chromatin which are bound by a member of the basal transcriptional machinery; and (d) comparing the promoter regions identified in steps (b) and (c) to determine the ratio between (i) the number of promoter regions bound by both the candidate global transcriptional regulator and the member of the basal transcriptional machinery; and (ii) the number of promoter regions bound by the member of the basal transcriptional machinery wherein a transcriptional regulator is a global transcriptional regulator when the ratio is greater than 0.2.

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In a preferred embodiment of the methods described above, steps (b) and (c) are performed using a DNA microarray. In a specific embodiment, the DNA microarray comprises (i) at least 10,000 experimental spots, each experimental spot comprising an experimental DNA, each experimental DNA comprising a promoter region from a human gene in the subset; and (ii) at least 100 control spots, each control spot comprising a control DNA, each control DNA comprising a non-promoter region. Any type of microarray or array may be used.

In one embodiment of the methods described above, the member of the transcriptional machinery is an RNA polymerase, such as RNA polymerase II, a TATA-binding protein, or any other component of TFIID, including, for example, the TAFs (e.g. TAF250, TAF150, TAF135, TAF95, TAF80, TAF55, TAF31, TAF28, and TAF20).

Another aspect of the invention provides methods of identifying regulatory networks, or pathways, in a cell. The methods provided by the invention allow the identification of the regulatory motifs, such as those shown in Figure 2B. A regulatory pathway can include, for example, a pathway that controls a cellular function under a

specific condition. A regulatory pathway controls a cellular function by, for example, altering the activity of a system component or the activity of a biochemical, gene expression or other type of pathway. Alterations in activity include, for example, inducing a change in the expression, activity, or physical interactions of a pathway component under a specific condition. Specific examples of regulatory pathways include a pathway that activates a cellular function in response to an environmental stimulus of a biochemical system, such as the inhibition of cell differentiation in response to the presence of a cell growth signal and the activation of galactose import and catalysis in response to the presence of galactose and the absence of repressing sugars. The term "component" when used in reference to a network or pathway is intended to mean a molecular constituent of the biochemical system, network or pathway, such as, for example, a polypeptide, nucleic acid, other macromolecule or other biological molecule.

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In one aspect, the invention provides a method of identifying a transcriptional regulatory network in a cell, the method comprising determining if a transcriptional regulator regulates additional transcriptional regulators in the cell, such as by using any of the methods described herein, wherein a transcriptional regulatory network is identified if at least one additional transcriptional regulator is regulated by the transcriptional regulator.:

Another aspect of the invention provides a method of identifying a transcriptional regulatory network in a cell, the method comprising determining if a transcriptional regulator regulates (i) its own promoter; or (ii) a promoter from a plurality of transcriptional regulators; such as by using any of the methods described herein, wherein the experimental DNA comprises (a) a promoter from the transcriptional regulator; and (b) promoters from the plurality of transcriptional regulators; wherein a transcriptional regulatory network is identified if the transcriptional regulator regulates itself or if it regulates at least one of the plurality of transcriptional regulators.

Yet another aspect of the invention provides a method of identifying

transcriptional regulatory networks in a cell, the method comprising (a) determining, by repeating one of the methods described herein for each of a plurality of transcriptional regulators, the genes in a subset which are regulated by each of the plurality of transcriptional regulators, wherein the experimental DNA comprises promoter regions for each of the plurality of transcriptional regulators; (b) determining if any one of the plurality of transcriptional regulators are regulated by at least one of the plurality of transcriptional regulators; wherein a transcriptional regulatory network is identified if any one of the plurality of transcriptional regulators is regulated by at least one of the plurality of transcriptional regulators.

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Specific embodiments of the methods for identifying regulatory networks described herein further comprise determining if any of the genes regulated by one of the plurality of transcriptional regulators is also a target of any of the other transcriptional regulators

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The invention further provides algorithms for the identification of regulatory motifs, which may be used in conjuction with any of the methods provided herein, such as the methods for identifying the genes regulated by a transcriptional regulator. In a specific embodiment, two data matrices are created. The overall matrix D consists of binary entries Dij, where a 1 indicates binding of regulator j to intergenic region i, a 0 indicates no binding event. The regulator matrix R is a subset of D, containing only the rows corresponding to the intergenic region assigned to each regulator, in the same order as the columns of regulators. The analyses may be performed using Matlab® software. The algorithms to find each motif are described as follows:

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Autoregulatory motif: Find each non-zero entry on the diagonal of R.

Feedforward loop: For each master regulator (column of R), find non-zero entries, which correspond to regulators bound. For each master regulator / secondary regulator pair, find all rows in D bound by both regulators.

Multi-component loop: For each regulator (column of R), find the regulators to

which it binds. For each of these, find the regulators it binds. If any of these are the original regulator, you have a multi-component loop of two. For all others, find regulators to which they bind. If any of these are the original, you have a multicomponent loop of three. Repeat to find larger loops.

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Single input module: Find the intergenic regions bound by only one regulator. That is, take the subset of rows of D such that the sum of each row is 1. Then for each regulator (column), find non-zero entries. Each set (greater than three intergenic regions) is a SIM.

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Multi-input module: Find the intergenic regions bound by more than one regulator. That is, take the subset of rows of D such that the sum of each row is greater than 1. Then, for each row, find any other row bound by the same regulators. The collection of rows bound by the same regulators correspond to a MIM. Once a row is assigned to a MIM, remove it from further analysis.

Regulator chain: For each regulator (column of R), use a recursive algorithm to find chains of all lengths. That is, for each regulator whose promoter is bound by the regulator before it in the chain, find the regulator promoters to which it binds. Repeat until the chain ends. There are three possible ways to end a chain: a regulator that does not bind to the promoter of any other regulator, a regulator that binds to its own promoter, or one that binds to the promoter of another regulator earlier in the chain.

In one preferred embodiment of any of the methods described herein such as the methods for identifying regulatory networks, the experimental DNA in the microarray comprises promoter regions from additional transcriptional regulators or from genes suspected to encode transcriptional regulators. Such microarray enables one skilled in the art to identify the components of a regulatory pathway. For example, starting with one transcriptional regulator, a subset of the genes it regulates are identified using any method, such as those described herein. If one identified gene is itself a second transcriptional regulator or is suspected to encode a transcriptional regulator, then the subset of genes the second transcriptional regulator regulates is identified, and so on.

Furthermore, the subset of genes that the first and second transcriptional regulators regulate can be compared to determine of any genes are found in both subsets. If so, then a feed-forward motif, a unit of a regulatory network, has been identified. Likewise, if the second transcriptional regulator is found to regulate the first one, then a feedback loop has been identified.

4. Development of a Therapeutic to Treat or Prevent Disorders

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One aspect of the invention provides methods of identifying targets for the development of therapeutics. One aspect of the invention provides a method of identifying at least one target gene for the development of a therapeutic to treat or prevent a disorder in a subject, wherein at least one form of the disorder is caused by an altered activity in a transcriptional regulator or in a suspected transcriptional regulator, the method comprising (a) identifying the genes regulated by the transcriptional regulator in a cell; (b) determining if the transcriptional regulator is a broad-acting transcriptional regulator or a narrow-acting transcriptional regulator, wherein if the transcriptional regulator is a broad acting transcriptional regulator then the transcriptional regulator is a target gene for the development of a therapeutic, and wherein if the transcriptional regulator is a narrow acting transcriptional regulator then (i) determining if at least one gene regulated by the transcriptional regulator is likely causative in the disorder, wherein a gene that is likely causative in the disorder is a target gene for the development of a therapeutic; and (ii) reiterating steps (a) and (b) for at least one gene that is regulated by the transcriptional regulator in the cell and that either (1) encodes a transcriptional regulator or (2) is suspected to encode a transcriptional regulator, with the modification that the transcriptional regulator of steps (a) and (b) is said gene, thereby identifying at least one target gene for the development of a therapeutic to treat or prevent a disorder in the subject.

In some embodiments of the methods for identifying a target gene for the development of a therapeutic, the genes regulated by the transcriptional regulator in the cell are identified using chromosome-wide location analysis, analysis of mRNA transcripts in a cell that expresses the transcriptional regulator, or by using any of the methods provided herein for the identification of the genes that are regulated by a

transcriptional regulator. Some methods may comprise the use of DNA microarray or DNA arrays, such as those described in Gabrielson et al., Obesity Research, 8(5), 374-384 (2000).

In some embodiments of the methods described herein for identifying a target gene for the development of a therapeutic, the transcriptional regulator is a master regulatory gene. In specific embodiments, the master regulatory gene is SOX1-18, OCT6, PAX3, Myocardin, GATA1-6, TCF1/HNF1A, HNF4A, HNF6, NGN3, C/EBP, FOXA1-3, IPF1, GATA, HNF3, NKX2.1, CDX, FTF/NR5A2, C/EBPbeta, SCL1, SKIN1, or a member of the neurogenin, LK, LMO, SOX, OCT, PAX, GATA or MyoD family of transcription factors.

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In some embodiments of the methods described herein, the transcriptional regulator is PAX3, EGR-1, EGR-2, OCT6, a SOX family member, a GATA family member, a PAX family member, an OCT family member, RFX5, WHN, GATA1, VDR, CRX, CBP, MeCP2, AML1, p53, PLZF, PML, Rb, WT1, NR3C2, GCCR, PPARgamma, SIM1, HNF1alpha, HNF1beta, HNF4alpha, PDX1, MAFA, FOXA2, or NEUROD1.

A transcriptional regulator whose altered activity can lead to disease might be expressed in multiple, or all tissues of an organism, such that any of multiple cell types may be used in identifying a therapeutic. In some embodiments of the methods described herein for identifying a target gene for the development of a therapeutic, the cell is derived from a tissue whose function is impaired in the disorder. For example, a pancreatic cell may be used for diabetes, a cardiac muscle cells for myocardial infarction, or neurons for Alzheimer's disease.

In specific embodiments of the methods described herein for identifying a target gene for the development of a therapeutic, the broad acting gene regulates at least about 1%, 2% or more preferably at least about 2.5% of the genes in the cell, and the narrow acting gene regulates less than about 1%, 2% or 2.5% of the genes in the cell.

In specific embodiments of the methods described herein, a gene is suspected to encode a transcriptional regulator if it shares at least about 30%, 40% or 50% amino acid sequence identity within at least the DNA binding domain of a transcriptional regulator. DNA binding domains and methods of performing nucleic acids and polypeptide sequence alignments are well-known in the art. Optimal alignment of sequences for comparison may be conducted by the local homology algorithm of Smith and Waterman, Adv. Appl. Math. 2: 482 (1981); by the homology alignment algorithm of Needleman and Wunsch, J. Mol Biol. 48: 443 (1970); by the search for similarity method of Pearson and Lipman, Proc. Natl. Acad. Sci. 8: 2444 (1988); by computerized implementations of these algorithms, including, but not limited to: CLUSTAL in the PC/Gene program by Intelligenetics, Mountain View, Calif., GAP, BESTFIT, BLAST, FASTA, and TFASTA in the Wisconsin Genetics Software Package, Genetics Computer Group (GCG), 7 Science Dr., Madison, Wis., USA; the CLUSTAL program is well described by Higgins and Sharp, Gene, 73: 237-244, 1988; Higgins and Sharp, CABIOS:11-13, 1989; Corpet, et al., Nucleic Acids Research, 16:881-90,1988; Huang, et al., Computer Applications in the Biosciences 8:1-7,1992; and Pearson, et al., Methods in Molecular Biology 24:7-331,1994.

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In some specific embodiments of the methods described herein for identifying a target gene for the development of a therapeutic, the gene regulated by the transcriptional regulator is said to be likely causative of the disorder if a mutation in said gene results in at least one phenotype or symptom associated with the disorder. In another specific embodiment, the gene regulated by the transcriptional regulator is said to be likely causative of the disorder when the gene encodes an enzyme or signaling molecule which functions in a pathway that is impaired in the disorder. For example, if the disease is type II diabetes, a disorder characterized by hyperglycemia, then a gene regulated by the transcriptional regulator which encodes a sugar transporter, an enzyme involved in catalyzing a step of glycolysis or gluconeogenesis, or a gene which regulates insulin production, secretion or signaling is said to be likely causative or the disorder. In another specific embodiment, the gene regulated by the transcriptional regulator is said to be likely causative of the disorder if a mutant allele of the gene is genetically linked to a "susceptibility locus" for at least one form of the disease. A

"susceptibility locus" for a particular disease is a sequence or gene locus implicated in the initiation or progression of the disease. The susceptibility locus can be, for example, a gene or a microsatellite repeat, as identified by a microsatellite marker, or can be identified by a defined single nucleotide polymorphism. Generally, susceptibility genes implicated in specific diseases and their loci can be found in scientific publications, but may also be determined experimentally.

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In some embodiments of the methods described herein for identifying a target gene for the development of a therapeutic, the altered activity in the transcriptional regulator comprises at least one of the following: (a) an alteration in the binding affinity of the transcriptional regulator to DNA; (b) an alteration in the ability of the transcriptional regulator to bind to RNA polymerase, to an RNA polymerase holoenzyme, or to a second transcriptional regulator; (c) an alteration in the binding affinity of the transcriptional regulator to a ligand; (d) an alteration in expression level or expression pattern of the transcriptional regulator; or (e) an alteration in an ability of the transcriptional regulator to form homomultimers or heteromultimers.

In some embodiments of the methods described herein, the cell comprises a mutant form of the transcriptional regulator. A preferred mutant form of the transcriptional regulator is one that causes the disease to which the therapeutic is sought. Such embodiments are particularly preferred when a mutant transcriptional regulator which causes at least one form of the disease has an altered target specificity and thus the genes it regulates, or the extent to which it regulates their transcription, is altered when compared to the non-mutant form of the transcriptional regulator. Such embodiments may allow the identification of therapeutic targets which might not have been identified if a wild-type form of the transcriptional regulator had been used. Mutations in the DNA binding domain, for example, may alter the target specificity of a transcriptional regulator by altering its affinity for various DNA binding sequences.

It is well-known to one skilled in the art that mutations in a transcriptional regulator may result in a hypomorphic, hypermorphic or neomorphic phenotype.

Mutations may generally reduce the activity of a transcriptional regulator, may

generally increase it activity, or may confer novel properties, such as altering the range of targets or turning an activator into a repressor or vice versa. In any methods described herein, and in particular those for identifying the therapeutics, a cell expressing a transcriptional regulator having any of these changes in activity may be used.

The methods described herein may be applied to any disorder for which a transcriptional regulator has been implicated. Examples of diseases and transcriptional regulators which cause them may be found in the scientific and medical literature by one skilled in the art, including in Medical Genetics, L.V. Jorde et al., Elsevier Science 2003, and Principles of Internal Medicine, 15th edition, ed by Braunwald et al., McGraw-Hill, 2001; American Medical Association Complete Medical Encyclopedia (Random House, Incorporated, 2003); and The Mosby Medical Encyclopedia, ed by Glanze (Plume, 1991). In some embodiments, the disorder is characterized by impaired function of at least one of the following: brain, spinal cord, heart, arteries, esophagus, stomach, small intestine, large intestine, liver, pancreas, lungs, kidney, urinary tract, ovaries, breasts, uterus, testis, penis, colon, prostate, bone, muscle, cartilage, thyroid gland, adrenal gland, pituitary, bone marrow, blood, thymus, spleen, lymph nodes, skin, eye, ear, nose, teeth or tongue.

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In some embodiments of the methods described herein for identifying a target gene for the development of a therapeutic, the subject is a mammal. In preferred embodiments, the subject is a human. In some embodiments of the methods described herein for identifying a target gene for the development of a therapeutic, the therapeutic comprises a small molecule drug, an antisense nucleic acid, an antibody, a peptide, a ligand, a fatty acid, a hormone or a metabolite.

Antisense nucleic acids acting by RNAi include oligonucleotides which specifically hybridize (e.g., bind) under cellular conditions with a gene sequence, such as at the cellular mRNA and/or genomic DNA level, so as to inhibit expression of that gene, e.g., by inhibiting transcription and/or translation. The binding may be by conventional base pair complementarily, or, for example, in the case of binding to DNA

duplexes, through specific interactions in the major groove of the double helix.

Preferred antisense nucleic acid comprise siRNA, shRNAs, or any other form of double stranded RNA molecule. Antisense nucleic acids may be chemically modified, such as to increase their in vivo stability.

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RNAi is a process of sequence-specific post-transcriptional gene repression which can occur in eukaryotic cells. In general, this process involves degradation of an mRNA of a particular sequence induced by double-stranded RNA (dsRNA) that is homologous to that sequence. For example, the expression of a long dsRNA corresponding to the sequence of a particular single-stranded mRNA (ss mRNA) will labilize that message, thereby "interfering" with expression of the corresponding gene. Accordingly, any selected gene may be repressed by introducing a dsRNA which corresponds to all or a substantial part of the mRNA for that gene. It appears that when a long dsRNA is expressed, it is initially processed by a ribonuclease III into shorter dsRNA oligonucleotides of in some instances as few as 21 to 22 base pairs in length. Furthermore, RNAi may be effected by introduction or expression of relatively short homologous dsRNAs. dsRNAs shorter than about 30 bases pairs are preferred to effect gene repression by RNAi (see Hunter et al. (1975) J Biol Chem 250: 409-17; Manche et al. (1992) Mol Cell Biol 12: 5239-48; Minks et al. (1979) J Biol Chem 254: 10180-3; and Elbashir et al. (2001) Nature 411: 494-8).

Antibodies include whole antibodies, e.g., of any isotype (IgG, IgA, IgM, IgE, etc.), and includes fragments thereof which are also specifically reactive with a vertebrate, e.g., mammalian, protein. Antibodies may be fragmented using conventional techniques and the fragments screened for utility in the same manner as described above for whole antibodies. Thus, the term includes segments of proteolytically-cleaved or recombinantly-prepared portions of an antibody molecule that are capable of selectively reacting with a certain protein. Non-limiting examples of such proteolytic and/or recombinant fragments include Fab, F(ab')2, Fab', Fv, and single chain antibodies (scFv) containing a V[L] and/or V[H] domain joined by a peptide linker. The scFv's may be covalently or non-covalently linked to form antibodies having two or more binding sites. The subject invention includes polyclonal, monoclonal,

humanized, or other purified preparations of antibodies and recombinant antibodies.

Peptidomimetic include compounds containing peptide-like structural elements that is capable of mimicking the biological action (s) of a natural parent polypeptide.

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Hormone include any one of a number of biochemical substances that are produced by a certain cell or tissue and that cause a specific biological change or activity to occur in another cell or tissue located elsewhere in the body.

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Metabolites includes any substance produced by metabolism or by a metabolic process. "Metabolism", as used herein, refers to the various chemical reactions involved in the transformation of molecules or chemical compounds occurring in tissue and the cells therein.

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Ligands include any substance which binds to a receptor protein. A ligand of a transcriptional regulator protein is a substance which binds to the regulator protein, such as estrogen binding to a nuclear hormone receptor. In a preferred embodiment, ligand binding of to a transcriptional regulator occurs with high affinity. The term ligand refers to substances including, but not limited to, a natural ligand, whether isolated and/or purified, synthetic, and/or recombinant, a homolog of a natural ligand (e.g., from another mammal). The term ligand encompasses substances which are inhibitors or promoters of receptor activity, as well as substances which selectively bind receptors, but lack inhibitor or promoter activity.

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Some aspects of the invention relate to the diagnosis of disease states. A "transcriptional fingerprint", or listing of the genes, and optionally to what extent, that are regulated by given a transcriptional regulator can be generated from healthy individuals and from those afflicted with a disorder. Comparison of the fingerprints between the two groups may define genes which are specific to one of the two groups, and thus serve as diagnostic for the risk that a patient is at risk, or is afflicted, with the disorder. In one embodiment, the transcriptional fingerprint of HNF4a is used to diagnose type II diabetes. A biopsy of a subject's liver or pancreas may provide the

cells for such analysis.

In specific embodiments, the transcriptional fingerprint disease diagnosis analysis is applied to transcriptional regulators which are causative in a particular disease to diagnose the disease. This approach may be coupled to allelic genotyping of the transcriptional regulator gene in the subject. For example, genotyping of a subject's HNF4a may uncover a novel allele. By using "transcriptional fingerprint" of HNF4a in tissue from that patient, one skilled in the art may determine what effect that mutation has in HNF4a activity and thus diagnose type II diabetes.

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5. Methods of Preventing/Treating Disease through Regulation of HNFs

Some aspects of the invention provide methods of treating or preventing disease by regulating transcriptional regulator activity, particularly that of the HNF family member. The invention provides a method of treating or preventing type II diabetes in a subject, comprising administering to the subject a therapeutically effective amount of an agent that increases the global transcriptional activity of HNF4alpha. U.S. Patent No. 5,849,485 describes methods and assays for the isolation of modulators of HNF-4a activity, hereby incorporated by reference.

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The invention also provides a method of treating or preventing a disorder associated with low transcriptional activity of HNF4alpha in a subject, comprising administering to the subject a therapeutically effective amount of an agent that increases the global transcriptional activity of HNF4alpha. In a related aspect, the invention provides a method of treating or preventing a disorder associated with high transcriptional activity of HNF4alpha in a subject, comprising administering to the subject a therapeutically effective amount of an agent that decreases the global transcriptional activity of HNF4alpha.

Yet another related aspect of the invention provides a method of increasing the global transcriptional activity in a liver or a pancreatic cell comprising contacting the cell with an agent which increases the global transcriptional activity of HNF4alpha. Similarly, the invention provides a method of decreasing the global transcriptional

activity in a liver or a pancreatic cell comprising contacting the cell with an agent which decreases the global transcriptional activity of HNF4alpha.

Applicants have identified genes that are transcriptionally regulated by HNF-1a, HNF4a and HNF6 in hepatocytes and pancreatic cells. Accordingly, the invention provides methods of regulating the expression level of any of these genes in a cell or in a subject by contacting the cell or administering to the subject and agent which modulates the expression level or transcriptional regulatory activity of HNF transcription factors.

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The invention provides a method of regulating the expression level of any one of the genes in Figure 13 in a hepatocyte, the method comprising contacting the cell with an agent which regulates the transcriptional activity of HNF1alpha. Similarly, the invention also provides a method of regulating the expression level of any one of the genes in Figure 14 in a pancreatic cell, the method comprising contacting the cell with an agent which regulates the transcriptional activity of HNF1alpha.

The invention also provides a method of regulating the expression level of any one of the genes in Figure 16 in a hepatocyte, the method comprising contacting the cell with an agent which regulates the transcriptional activity of HNF6. Similarly, the invention provides a method of regulating the expression level of any one of the genes in Figure 17 in a pancreatic cell, the method comprising contacting the cell with an agent which regulates the transcriptional activity of HNF6.

The invention additionally provides a method of regulating the expression level of any one of the genes in Figure 18 in a hepatocyte, the method comprising contacting the cell with an agent which regulates the transcriptional activity of HNF4alpha. Similarly, the invention provides a method of regulating the expression level of any one of the genes in Figure 19 in a pancreatic cell, the method comprising contacting the cell with an agent which regulates the transcriptional activity of HNF4alpha.

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Agents which modulate the transcriptional activity of HNF-4a, or any other HNF family member, may be identified by screening compounds for their ability to

increase the expression level, the DNA binding activity or the transcriptional promoting activity of HNF4a. One assay format which can be used employs two genetic constructs. One is typically a plasmid that continuously expresses the transcriptional regulator of interest when transfected into an appropriate cell line. CV-1 cells are most often used. The second is a plasmid which expresses a reporter, e.g., luciferase under control of the transcriptional regulator. For example, if a compound which acts as a ligand for HNF-4 is to be evaluated, one of the plasmids would be a construct that results in expression of the HNF-4 receptor in an appropriate cell line, e.g., the CV-1 cells. The second would possess a promoter linked to the luciferase gene in which an HNF-4 response element is inserted. If the compound to be tested is an agonist for the HNF-4 receptor, the ligand will complex with the receptor and the resulting complex binds the response element and initiates transcription of the luciferase gene. In time the cells are lysed and a substrate for luciferase added. The resulting chemiluminescence is measured photometrically. Dose response curves are obtained and can be compared to the activity of known ligands. Other reporters than luciferase can be used including CAT and other enzymes.

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Viral constructs can be used to introduce the gene for the receptor and the reporter. An usual viral vector is an adenovirus. For further details concerning this preferred assay, see U.S. Pat. No. 4,981,784 issued Jan. 1, 1991 hereby incorporated by reference, and Evans et al., WO88/03168 published on 5 May 1988, also incorporated by reference.

HNF-4a antagonists can be identified using this same basic "agonist" assay. A fixed amount of an antagonist is added to the cells with varying amounts of test compound to generate a dose response curve. If the compound is an antagonist, expression of luciferase is suppressed.

Additional methods for the isolation of agonists and antagonist of HNF transcription factors are described in U.S. Patent Nos. 6,187,533 and 5,620,887.

Additional U.S. patents describing methods to identify agents that modulate the activity of transcription factors include 5,804,374, and 5,298,429, and U.S. Patent Publication

Nos. 2004/0033942A1 2003/0077664, 2003/0215829 and 2003/0039980. Any of the methods described herein may be easily adapted to identify agonists or antagonists of any one of the HNF transcriptional factors. U.S. Patent No. 6,303,653 describes modulators of HNF-4 activity.

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Agonists and antagonists of HNF4a can also be designed based on the known crystal structure of HNF4a complexed with an endogenous fatty acid ligand (Dhe-Paganon, J. Biol. Chem. 277(41), 37973-37976). U.S. Patent Publication No. 2002/0072587 describes methods of identifying agonists of an estrogen receptor, a nuclear receptor like the HNF proteins, based on its crystal structure. Such methods may easily be applied to HNF-1a, HNF-4a and HNF6 by one skilled in the art. Additional examples of rational drug design based on the structure of a protein may be found in U.S. Patent or Publication Nos. 6,236,946, 6,684,162, 2004/0014153, 2003/0124699, 20030077628, 2002/0151028, 2002/0072587 and 2003/0211588.

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6. Therapeutics

In one aspect, the invention provides methods of treating disease in a subject comprising the administration of a composition comprising a therapeutic agent. "Therapeutic agent" or "therapeutic" refers to an agent capable of having a desired biological effect on a host. Chemotherapeutic and genotoxic agents are examples of therapeutic agents that are generally known to be chemical in origin, as opposed to biological, or cause a therapeutic effect by a particular mechanism of action, respectively. Examples of therapeutic agents of biological origin include growth factors, hormones, and cytokines. A variety of therapeutic agents are known in the art and may be identified by their effects. Certain therapeutic agents are capable of regulating cell proliferation and differentiation. Examples include chemotherapeutic nucleotides, drugs, hormones, non-specific (non-antibody) proteins, oligonucleotides (e.g., antisense oligonucleotides that bind to a target nucleic acid sequence (e.g., mRNA sequence)), peptides, and peptidomimetics.

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In one embodiment, the compositions are pharmaceutical compositions.

Pharmaceutical compositions for use in accordance with the present invention may be

formulated in conventional manner using one or more physiologically acceptable carriers or excipients. Thus, the compounds and their physiologically acceptable salts and solvates may be formulated for administration by, for example, by aerosol, intravenous, oral or topical route. The administration may comprise intralesional, intraperitoneal, subcutaneous, intramuscular or intravenous injection; infusion; liposome-mediated delivery; topical, intrathecal, gingival pocket, per rectum, intrabronchial, nasal, transmucosal, intestinal, oral, ocular or otic delivery.

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An exemplary composition of the invention comprises an compound capable of modulating the expression or activity of a transcriptional regulator with a delivery system, such as a liposome system, and optionally including an acceptable excipient. In a preferred embodiment, the composition is formulated for injection.

Techniques and formulations generally may be found in Remmington's Pharmaceutical Sciences, Meade Publishing Co., Easton, PA. For systemic administration, injection is preferred, including intramuscular, intravenous, intraperitoneal, and subcutaneous. For injection, the compounds of the invention can be formulated in liquid solutions, preferably in physiologically compatible buffers such as Hank's solution or Ringer's solution. In addition, the compounds may be formulated in solid form and redissolved or suspended immediately prior to use. Lyophilized forms are also included.

For oral administration, the pharmaceutical compositions may take the form of, for example, tablets or capsules prepared by conventional means with pharmaceutically acceptable excipients such as binding agents (e.g., pregelatinised maize starch, polyvinylpyrrolidone or hydroxypropyl methylcellulose); fillers (e.g., lactose, microcrystalline cellulose or calcium hydrogen phosphate); lubricants (e.g., magnesium stearate, talc or silica); disintegrants (e.g., potato starch or sodium starch glycolate); or wetting agents (e.g., sodium lauryl sulphate). The tablets may be coated by methods well known in the art. Liquid preparations for oral administration may take the form of, for example, solutions, syrups or suspensions, or they may be presented as a dry product for constitution with water or other suitable vehicle before use. Such liquid

preparations may be prepared by conventional means with pharmaceutically acceptable additives such as suspending agents (e.g., sorbitol syrup, cellulose derivatives or hydrogenated edible fats); emulsifying agents (e.g., lecithin or acacia); non-aqueous vehicles (e.g., ationd oil, oily esters, ethyl alcohol or fractionated vegetable oils); and preservatives (e.g., methyl or propyl-p-hydroxybenzoates or sorbic acid). The preparations may also contain buffer salts, flavoring, coloring and sweetening agents as appropriate.

Preparations for oral administration may be suitably formulated to give controlled release of the active compound. For buccal administration the compositions may take the form of tablets or lozenges formulated in conventional manner. For administration by inhalation, the compounds for use according to the present invention are conveniently delivered in the form of an aerosol spray presentation from pressurized packs or a nebuliser, with the use of a suitable propellant, e.g., dichlorodifluoromethane, trichlorofluoromethane, dichlorotetrafluoroethane, carbon dioxide or other suitable gas. In the case of a pressurized aerosol the dosage unit may be determined by providing a valve to deliver a metered amount. Capsules and cartridges of e.g., gelatin for use in an inhaler or insufflator may be formulated containing a powder mix of the compound and a suitable powder base such as lactose or starch.

The compounds may be formulated for parenteral administration by injection, e.g., by bolus injection or continuous infusion. Formulations for injection may be presented in unit dosage form, e.g., in ampoules or in multi-dose containers, with an added preservative. The compositions may take such forms as suspensions, solutions or emulsions in oily or aqueous vehicles, and may contain formulatory agents such as suspending, stabilizing and/or dispersing agents. Alternatively, the active ingredient may be in powder form for constitution with a suitable vehicle, e.g., sterile pyrogen-free water, before use.

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The compounds may also be formulated in rectal compositions such as suppositories or retention enemas, e.g., containing conventional suppository bases such

as cocoa butter or other glycerides.

WO 2005/054461

In addition to the formulations described previously, the compounds may also be formulated as a depot preparation. Such long acting formulations may be administered by implantation (for example subcutaneously or intramuscularly) or by intramuscular injection. Thus, for example, the compounds may be formulated with suitable polymeric or hydrophobic materials (for example as an emulsion in an acceptable oil) or ion exchange resins, or as sparingly soluble derivatives, for example, as a sparingly soluble salt.

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Systemic administration can also be by transmucosal or transdermal means. For transmucosal or transdermal administration, penetrants appropriate to the barrier to be permeated are used in the formulation. Such penetrants are generally known in the art, and include, for example, for transmucosal administration bile salts and fusidic acid derivatives. in addition, detergents may be used to facilitate permeation. Transmucosal administration may be through nasal sprays or using suppositories. For topical administration, the oligomers of the invention are formulated into ointments, salves, gels, or creams as generally known in the art. A wash solution can be used locally to treat an injury or inflammation to accelerate healing.

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The compositions may, if desired, be presented in a pack or dispenser device which may contain one or more unit dosage forms containing the active ingredient.

The pack may for example comprise metal or plastic foil, such as a blister pack. The pack or dispenser device may be accompanied by instructions for administration.

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For therapies involving the administration of nucleic acids, the oligomers of the invention can be formulated for a variety of modes of administration, including systemic and topical or localized administration. Techniques and formulations generally may be found in Remmington's Pharmaceutical Sciences, Meade Publishing Co., Easton, PA. For systemic administration, injection is preferred, including intramuscular, intravenous, intraperitoneal, intranodal, and subcutaneous for injection, the oligomers of the invention can be formulated in liquid solutions, preferably in

physiologically compatible buffers such as Hank's solution or Ringer's solution. In addition, the oligomers may be formulated in solid form and redissolved or suspended immediately prior to use. Lyophilized forms are also included.

Systemic administration can also be by transmucosal or transdermal means, or the compounds can be administered orally. For transmucosal or transdermal administration, penetrants appropriate to the barrier to be permeated are used in the formulation. Such penetrants are generally known in the art, and include, for example, for transmucosal administration bile salts and fusidic acid derivatives. In addition, detergents may be used to facilitate permeation. Transmucosal administration may be through nasal sprays or using suppositories. For oral administration, the oligomers are formulated into conventional oral administration forms such as capsules, tablets, and tonics. For topical administration, oligomers may be formulated into ointments, salves, gels, or creams as generally known in the art.

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Toxicity and therapeutic efficacy of the agents and compositions of the present invention can be determined by standard pharmaceutical procedures in cell cultures or experimental animals, e.g., for determining the LD50 (the dose lethal to 50% of the population) and the ED50 (the dose therapeutically effective in 50% of the population). The dose ratio between toxic and therapeutic effects is the therapeutic index and it can be expressed as the ratio LD50/ED50. Compounds which exhibit large therapeutic induces are preferred. While compounds that exhibit toxic side effects may be used, care should be taken to design a delivery system that targets such compounds to the site of affected tissue in order to minimize potential damage to uninfected cells and, thereby, reduce side effects.

The data obtained from the cell culture assays and animal studies can be used in formulating a range of dosage for use in humans. The dosage of such compounds lies preferably within a range of circulating concentrations that include the ED50 with little or no toxicity. The dosage may vary within this range depending upon the dosage form employed and the route of administration utilized. For any compound used in the method of the invention, the therapeutically effective dose can be estimated initially

from cell culture assays. A dose may be formulated in animal models to achieve a circulating plasma concentration range that includes the IC50 (i.e., the concentration of the test compound which achieves a half-maximal inhibition of symptoms) as determined in cell culture. Such information can be used to more accurately determine useful doses in humans. Levels in plasma may be measured, for example, by high performance liquid chromatography.

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In one embodiment of the methods described herein, the effective amount of the agent is between about 1mg and about 50mg per kg body weight of the subject. In one embodiment, the effective amount of the agent is between about 2mg and about 40mg per kg body weight of the subject. In one embodiment, the effective amount of the agent is between about 3mg and about 30mg per kg body weight of the subject. In one embodiment, the effective amount of the agent is between about 4mg and about 20mg per kg body weight of the subject. In one embodiment, the effective amount of the agent is between about 5mg and about 10mg per kg body weight of the subject.

In one embodiment of the methods described herein, the agent is administered at least once per day. In one embodiment, the agent is administered daily. In one embodiment, the agent is administered every other day. In one embodiment, the agent is administered every 6 to 8 days. In one embodiment, the agent is administered weekly.

As for the amount of the compound and/or agent for administration to the subject, one skilled in the art would know how to determine the appropriate amount. As used herein, a dose or amount would be one in sufficient quantities to either inhibit the disorder, treat the disorder, treat the subject or prevent the subject from becoming afflicted with the disorder. This amount may be considered an effective amount. A person of ordinary skill in the art can perform simple titration experiments to determine what amount is required to treat the subject. The dose of the composition of the invention will vary depending on the subject and upon the particular route of administration used. In one embodiment, the dosage can range from about 0.1 to about 100,000 ug/kg body weight of the subject. Based upon the composition, the dose can be

delivered continuously, such as by continuous pump, or at periodic intervals. For example, on one or more separate occasions. Desired time intervals of multiple doses of a particular composition can be determined without undue experimentation by one skilled in the art.

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The effective amount may be based upon, among other things, the size of the compound, the biodegradability of the compound, the bioactivity of the compound and the bioavailability of the compound. If the compound does not degrade quickly, is bioavailable and highly active, a smaller amount will be required to be effective. The effective amount will be known to one of skill in the art; it will also be dependent upon the form of the compound, the size of the compound and the bioactivity of the compound. One of skill in the art could routinely perform empirical activity tests for a compound to determine the bioactivity in bioassays and thus determine the effective amount. In one embodiment of the above methods, the effective amount of the compound comprises from about 1.0 ng/kg to about 100 mg/kg body weight of the subject. In another embodiment of the above methods, the effective amount of the compound comprises from about 100 ng/kg to about 50 mg/kg body weight of the subject. In another embodiment of the above methods, the effective amount of the compound comprises from about 1 ug/kg to about 10 mg/kg body weight of the subject. In another embodiment of the above methods, the effective amount of the compound comprises from about 100 ug/kg to about 1 mg/kg body weight of the subject.

As for when the compound, compositions and/or agent is to be administered, one skilled in the art can determine when to administer such compound and/or agent. The administration may be constant for a certain period of time or periodic and at specific intervals. The compound may be delivered hourly, daily, weekly, monthly, yearly (e.g. in a time release form) or as a one time delivery. The delivery may be continuous delivery for a period of time, e.g. intravenous delivery. In one embodiment of the methods described herein, the agent is administered at least once per day. In one embodiment of the methods described herein, the agent is administered daily. In one embodiment of the methods described herein, the agent is administered every other day. In one embodiment of the methods described herein, the agent is administered every other day.

to 8 days. In one embodiment of the methods described herein, the agent is administered weekly.

5 EXEMPLIFICATION

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The invention now being generally described, it will be more readily understood by reference to the following examples, which are included merely for purposes of illustration of certain aspects and embodiments of the present invention, and are not intended to limit the invention, as one skilled in the art would recognize from the teachings hereinabove and the following examples, that other DNA microarrays, transcriptional regulators, cell types, antibodies, ChIP conditions, or data analysis methods, all without limitation, can be employed, without departing from the scope of the invention as claimed.

The practice of the present invention will employ, where appropriate and unless otherwise indicated, conventional techniques of cell biology, cell culture, molecular biology, transgenic biology, microbiology, virology, recombinant DNA, and immunology, which are within the skill of the art. Such techniques are described in the literature. See, for example, Molecular Cloning: A Laboratory Manual, 3rd Ed., ed. by Sambrook and Russell (Cold Spring Harbor Laboratory Press: 2001); the treatise, Methods In Enzymology (Academic Press, Inc., N.Y.); Using Antibodies, Second Edition by Harlow and Lane, Cold Spring Harbor Press, New York, 1999; Current Protocols in Cell Biology, ed. by Bonifacino, Dasso, Lippincott-Schwartz, Harford, and Yamada, John Wiley and Sons, Inc., New York, 1999; and PCR Protocols, ed. by Bartlett et al., Humana Press, 2003.

Various publications, patents, and patent publications are cited throughout this application the contents of which are incorporated herein by reference in their entirety.

30 Experimental procedures

The following procedures were followed in performing the experiments below:

Genome-scale Location Analysis

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The protocol described here was adapted from Ren 2001. Briefly, cells are fixed with 1% final concentration formaldehyde for 10-20 minutes at room temperature, harvested and rinsed with 1x PBS. The resultant cell pellet is sonicated, and DNA fragments that are crosslinked to a protein of interest are enriched by immunoprecipitation with a factor specific antibody. After reversal of the crosslinking, the enriched DNA is amplified using ligation-mediated PCR (LM-PCR), and then fluorescently labeled using high concentration Klenow polymerase and a dNTPfluorophore. A sample of DNA that has not been enriched by immunoprecipitation is subjected to LM-PCR and labeled with a different fluorophore. Both IP-enriched and unenriched pools of labeled DNA are hybridized to a single DNA microarray containing 13,000 human intergenic regions (see below for description of DNA microarray and binding site determination). For hepatocyte experiments, 2.5 x 107 hepatocytes were typically used per chromatin immunoprecipitation. These hepatocytes were isolated by standard liver perfusion techniques, immediately crosslinked with 1% formaldehyde solution, rinsed, and flash frozen. Islet preparations were treated with formaldehyde between 1 hour and 5 days after isolation from pancreata. A minimum of 30,000 viable islet equivalents (approximately 2x 10⁷ beta cells) were fixed and handled as described above. Typical islet purity for three experiments described here was >70% islets with >80% viability. HNF4a, HNF6, and RNA polymerase II produced high quality results with as few as 30,000 islet equivalents. HNF1a ChIP required significantly more material, typically 80,000 islets, to produce results with somewhat lower enrichment ratios than the results obtained with hepatocytes.

25 Human 13K DNA Microarray

It would be ideal to have a DNA microarray that contains the entire human genome sequence, but technical limitations and cost led applicants to select the most relevant portion of the genome for inclusion in this microarray. Because a significant percentage of transcriptional binding sites in proximal promoters are within 1 kb of transcription start sites, applicants designed primers to amplify these genomic regions for printing onto a promoter array. Applicants selected 15000 cDNAs from the NCBI RefSeq database, and mapped them to NCBI Build 22 (April 2001) of the human

genome using BLAST. Where multiple splice variants had been described, applicants used the most upstream site, and verified the 5'-end by alignment with the Database of Transcriptional Start Sites (http://elmo.ims.utokyo.ac.jp/dbtss/). Sequences to be amplified were extracted from the genomic region–750 bp to +250 bp relative to this transcriptional start site. To control for nonspecific binding, 9 amplified regions derived from long Arabidopsis open reading frames were included on the array. As a further negative control and for use in data normalization, applicants chose 158 ORF regions within long exons of human genes for amplification. To prepare the DNA content of the arrays, the program Primer3

(http://wwwgenome.wi.mit.edu/genome_software/other/ primer3.html) was used to design primers using the sequences described above. PCRs were performed on these primer set using standard conditions, except for the presence of 1 M betaine in all PCR reactions. Betaine was empirically observed to increase the success rate of the amplification reactions.

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Of the 13,000 PCR pairs, 70% gave a strong band of the appropriate size, as verified on 2% agarose gels. Applicants have noted, however, that PCR products undetectable by agarose EtBr gel analysis can give valid positive signals when concentrated and printed on the DNA arrays. PCR quality evaluations were performed on the BRIDNAsuite of programs from the Biotechnology Research Institute of the National Research Council of Canada (http://www.irb-bri.cnrc-nrc.gc.ca/).PCR products were recovered from the reaction mixture by ammonium acetate/isopropanol precipitation and resuspended into 3x SSC with 1.5 M betaine to minimize evaporation and improve spot quality. Applicants printed amplified products onto GAPS-coated glass slides (Corning) using a Cartesian PixSys 5500 arrayer. The quality of the arrays was determined on a batch-wise basis by hybridization with sequence neutral oligonucleotides covalently linked to Cy3 or Cy5, followed by calculation of usable percentage of spots, combined with direct visual inspection of the quality of the chip. The Hu13K array was remapped post-production using two independent methods. First, applicants performed electronic PCR on the primer sets against the August 2003 final release of the completed human genome. Second, applicants BLASTed the sequence used to extract primers for amplification against the August 2003 final release of the

human genome. The dataset downloadable from the supporting website reports the location of each arrayed promoter relative to the transcriptional start site.

Data Quality Control

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1. ChIP Hybridization Quality Control

The raw data generated from each array experiment was subjected to multiple levels of quality control. First, each scan was examined visually as it was being performed. Samples on microarrays with gross defects (e.g. scratches, smeared spots) were repeated whenever possible. Applicants also determined that no reliable signal was produced from control spots containing *Arabidopsis* DNA.

2. Binding Site Determination and Error Model

Scanned images were analyzed using GenePix (v3.1 or v4.0), to obtain background subtracted intensity values. Each spot is bound by both IP-enriched and unenriched DNA, which are labeled with different fluorophores. Consequently, each spot yields fluorescence intensity information in two channels, corresponding to immunoprecipitated DNA and genomic DNA. To account for background hybridization to slides, the median intensity of a set of control blank spots was subtracted for sitespecific transcription factors (e.g. HNF1a), and the median intensity for a set of control ORF spots was subtracted for broadly acting DNA binding proteins (e.g. RNA Pol II. HNF4a). To correct for different amounts of genomic and immunoprecipitated DNA hybridized to the microarray, the median intensity value of the IP-enriched DNA channel was divided by the median of the genomic DNA channel, and this normalization factor was applied to each intensity in the genomic DNA channel. Next, applicants calculated the log of the ratio of intensity in the IP-enriched channel to intensity in the genomic DNA channel for each intergenic region across the entire set of hybridization experiments. Adjusted intensity values for the IP-enriched channel were calculated from these ratios. A whole-chip error model (Hughes 2000; Lee 2002) was then used to calculate confidence values for each spot on each microarray, and to combine data for the replicates of each experiment to obtain a final average ratio and confidence for each promoter region. Genes were included in the set of 'bound' genes if the binding P-value in the error model was < 0.001 or enrichment was at least 2-fold

in the immunoprecipitation.

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Confirmation of Predicted Binding

The accuracy of genome-wide location data reported here has been assessed using several approaches.

1. Estimation of False Positive Rates Using Conventional ChIP Experiments

Conventional, independent ChIP experiments conducted in our laboratory at a gene specific level have confirmed over 100 binding interactions identified by location analysis data involving 6 different regulators (see http://web.wi.mit.edu/young/pancregulators). These results suggest that our empirical rate of false positives is at most 16%. This rate is somewhat higher than that found for a large scale survey of yeast transcription factors (Lee 2002), which probably reflects the greater complexity of the human genome. Figures 9 and 10 show typical verification ChIP experiments for HNF4a and HNF1a, respectively, in hepatocytes.

2. Comparison with Previous Literature

Applicants found no previous studies of the genomic targets of transcriptional regulators in primary human tissue. However, a large number of HNF1a and HNF4a targets have been identified in model organisms and human carcinoma (mostly hepatoma) cell lines; these targets are summarized in Figure 14. For example, genomescale location analysis identified 30 of the 68 hepatocyte genes which were both previously suggested to be targets of HNF4a, and included on the 13K DNA array. Similarly, genome-scale location analysis identified 21 of the 81 hepatocyte genes which were both previously suggested to be targets of HNF4a, and included on the 13K DNA array. Discrepancies between the targets reported here and targets reported in the literature may result from a number of factors, which include, but are not limited to: (1) the limitations of using a 1 kb promoter fragment to probe the binding of a transcription factor, (2) the stringency of our threshold criteria, (3) the differences between the regulatory network in model organisms and/or cell lines, and the regulatory network in primary human tissue, (4) differences between indirect technologies in the literature (i.e. gel-shift and transient transfections) and genome-scale location analysis, (5) tissue isolation effects, among others. A more comprehensive discussion can be found at

http://web.wi.mit.edu/young/pancregulators

Regulatory Motifs Derived from Binding Data

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In order to discover network motifs, two data matrices were created. The overall matrix D consists of binary entries Dij, where a 1 indicates binding of regulator j to intergenic region i, a 0 indicates no binding event. The regulator matrix R is a subset of D, containing only the rows corresponding to the intergenic region assigned to each regulator, in the same order as the columns of regulators. All analyses were performed in Matlab. The algorithms used to find each motif are described below. Autoregulatory motif: Find each non-zero entry on the diagonal of R. Feedforward loop: For each master regulator (column of R), find non-zero entries, which correspond to regulators bound. For each master regulator / secondary regulator pair, find all rows in D bound by both regulators. Multi-component loop: For each regulator (column of R), find the regulators to which it binds. For each of these, find the regulators it binds. If any of these are the original regulator, you have a multi-component loop of two. For all others, find regulators to which they bind. If any of these are the original, you have a multicomponent loop of three. Repeat to find larger loops. Single input module: Find the intergenic regions bound by only one regulator. That is, take the subset of rows of D such that the sum of each row is 1. Then for each regulator (column), find non-zero entries. Each set (greater than three intergenic regions) is a SIM. Multi-input module: Find the intergenic regions bound by more than one regulator. That is, take the subset of rows of D such that the sum of each row is greater than 1. Then, for each row, find any other row bound by the same regulators. The collection of rows bound by the same regulators correspond to a MIM. Once a row is assigned to a MIM, remove it from further analysis. Regulator chain: For each regulator (column of R), use a recursive algorithm to find chains of all lengths. That is, for each regulator whose promoter is bound by the regulator before it in the chain, find the regulator promoters to which it binds. Repeat until the chain ends. There are three possible ways to end a chain: a regulator that does not bind to the promoter of any other regulator, a regulator that binds to its own promoter, or one that binds to the promoter of another regulator earlier in the chain.

Example 1

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The liver and pancreas have long been the subject of studies to understand how organs develop and are regulated at the transcriptional level (8-12). The transcriptional regulators HNF1 α (a homeodomain protein), HNF4 α (a nuclear receptor) and HNF6 (a member of the onecut family) operate cooperatively in a connected network in the liver, but less in known about the structure of this regulatory network in human pancreatic islets. All three transcriptional regulators are required for normal function of liver and pancreatic islets (13-18). Mutations in HNF1 α and HNF4 α are the causes of the type 3 and type 1 forms of maturity-onset diabetes of the young (MODY3 and MODY1), a genetic disorder of the insulin-secreting pancreatic beta cells characterized by onset of diabetes mellitus before 25 years of age and an autosomal dominant pattern of inheritance (19).

Applicants hypothesized that genome-scale analysis of the pancreatic islet genes whose expression is regulated by these transcription factors in normal beta cells could provide insights into the molecular basis of the abnormal beta cell function that characterizes MODY. Applicants have identified the genes occupied by the transcription factors HNF1α, HNF4α, and HNF6 in pancreatic islets. The genes transcribed in each tissue were identified by determining the genomic occupancy of RNA polymerase Π. Applicants used this information to begin to map the transcriptional regulatory circuitry in these tissues.

Applicants first used genome-scale location analysis (20) to identify the promoters bound by HNF1α in human hepatocytes and pancreatic islets isolated from tissue donors (Fig 1A). For each tissue, HNF1α-DNA complexes were enriched by chromatin immunoprecipitation in three separate experiments. Applicants constructed a custom DNA microarray containing portions of promoter regions of 13,000 human genes (Hu13K array). Applicants targeted the region spanning 700 bp upstream and 200 bp downstream of transcription start sites for the genes whose start sites are best characterized based on National Center for Biotechnology Information annotation (20). Although many enhancers are present at more distant locations, most known

transcription factor binding site sequences occur within these start-site proximal regions of promoters.

The results of these genome location experiments revealed that HNF1 α is bound to at least 222 target genes in hepatocytes, representing 1.6% of the genes on the Hu13K array (Figure 11) (20). This result was verified with independent, conventional chromatin immunoprecipitation experiments, which suggest that the frequency of false positives in genome-scale location data with gene-specific regulators is no more than 16% when our threshold criteria were used (20). The genes applicants found to be occupied by HNF1 α in primary human hepatocytes encode products whose functions represent a significant cross-section of hepatocyte biochemistry. The results confirm that HNF1 α contributes to the transcriptional regulation of many of the central rate-limiting steps in gluconeogenesis and associated pathways. HNF1 α also binds to genes whose products are central to normal hepatic function, including carbohydrate synthesis and storage, lipid metabolism (synthesis of cholesterol and apolipoproteins), detoxification (synthesis of cytochrome P450s) and synthesis of serum proteins (albumin, complements and coagulation factors).

Applicants next identified HNF1 α target genes in human pancreatic islets (Figure 11) (20). HNF1 α occupied the promoter regions of 106 genes (0.8% of the Hu13K array promoters) in islets, 30% of which were also bound by HNF1 α in hepatocytes (Figure 1B). In islets, fewer chaperones and enzymes are bound by HNF1 α than in hepatocytes, and the receptors and signal transduction machinery regulated by HNF1 α vary between the two tissues.

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HNF1 a has been previously implicated in the regulation of many genes in hepatocytes and islets (13, 16, 20 [Figure 15]). The direct genome binding data reported here confirmed many, but not all, of these genes. The difference may be due, at least in part, to our stringent criteria for binding in the genome-scale data, which enhances our confidence in the direct target genes identified by location analysis, but likely underestimates the actual number of targets in vivo. Furthermore, although the

proximal promoter regions printed on the array contain a significant number of transcription factor binding sequences, many genes are also regulated by more distal promoter elements and enhancers that are not present on the Hu13K array.

Applicants also identified the promoters bound by HNF6 in human hepatocytes and pancreatic islets using genome-scale location analysis (Fig 1B; Figures 16 and 17) (20). HNF6 was bound to at least 222 genes in hepatocytes and 189 genes in pancreatic islets, representing 1.7% and 1.4% of the promoters on the array, respectively. Approximately half of the promoters occupied by HNF6 were common to the two tissues, and included a number of important cell cycle regulators such as CDK2 (20).

Genome-scale location analysis revealed surprising results for HNF4 α in hepatocytes and pancreatic islets (Fig 1B). The number of genes enriched in HNF4 α chromatin immunoprecipitations was much larger than observed with typical site-specific regulators. HNF4 α was bound to approximately 12% of the genes represented on the Hu13K DNA microarray in hepatocytes and 11% in pancreatic islets. No other transcription factor applicants have profiled in human cells has been observed to bind more than 2.5% of the promoter regions represented on the 13K array.

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Six independent lines of evidence indicate that the HNF4 α results are not due to poor antibody specificity or errors in the microarray analysis, and support the view that HNF4 α is associated with an unusually large number of promoters in hepatocytes and pancreatic islets (20). First, essentially identical results were obtained with two different antibodies that recognize different portions of HNF4 α . Second, Western blots showed that the HNF4 α antibodies are highly specific. Third, applicants verified binding at over 50 randomly selected targets of HNF4 α in hepatocytes by conventional gene-specific chromatin immunoprecipitation. Fourth, when antibodies against HNF4 α were used for ChIP in control experiments with Jurkat, U937, and BJT cells (which do not express HNF4 α), no more than 17 promoters were identified in each cell line by our criteria, which is well within the noise inherent in this system. Fifth, when pre-immune antibodies from rabbit and goat (the two different anti-HNF4 α antibodies came from rabbit and goat) were used in control experiments in hepatocytes, the

number of targets identified was within the noise. Finally, if the HNF4 α results are correct, then applicants would expect that the set of promoters bound by HNF4 α should be largely a subset of those bound by RNA polymerase II in each tissue; applicants found that this is the case (see below). Applicants conclude that HNF4 α is a widely acting transcription factor in these tissues, consistent with the observation that it is an unusually abundant, constitutively active transcription factor (11).

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Applicants next identified the genes represented on the Hu13K microarray that are actively transcribed in hepatocytes and pancreatic islets, so the fraction of actively transcribed genes that are bound by HNF4\alpha could be determined (Fig 2C). It is difficult to determine accurately the transcriptome of these tissues by profiling transcript levels with DNA microarrays. Transcript profiling requires a reference RNA population against which a tissue RNA population can be compared, and there are limitations to generating appropriate reference RNA. To circumvent this limitation, applicants exploited the fact that RNA polymerase II occupies the set of protein-coding genes that are actively transcribed in eukaryotic cells. Location analysis with RNA polymerase II antibodies can identify these actively transcribed genes (7, 21). Applicants found that 23% of the genes on the Hu13K array (2984 genes) were bound by RNA polymerase II in hepatocytes, and 19% (2426 genes) were bound by RNA polymerase II in islets (20). The sets of genes occupied by RNA polymerase II in hepatocytes and islets overlapped substantially (81% overlap, relative to islets), consistent with the relatedness of the two tissues (22). As expected, the majority of genes occupied by HNF4α in hepatocytes and pancreatic islets (80% and 73%, respectively) were also occupied by RNA polymerase II. Remarkably, of the genes occupied by RNA polymerase II, 42% (1262/2984) were bound by HNF4 α in hepatocytes and 43% (1047/2426) were bound by HNF4 α in islets (Fig 1C). By comparison, only 6% and 2% of RNA polymerase II enriched promoters were also bound by HNF1 α in hepatocytes and islets, respectively.

Previous studies indicate that HNF1α, HNF4α, and HNF6 are at the center of a network of transcription factors that cooperatively regulate numerous developmental and metabolic functions in hepatocytes and islets (9, 13, 15, 17). Our systematic

analysis of the direct in vivo targets of these factors significantly expands our understanding of the regulatory network in primary human tissues (Fig 2A). A comparison of the regulatory network in these two tissues reveals that HNF1 α , HNF4 α , and HNF6 occupy the promoters of genes encoding a large population of transcription factors and cofactors in the two tissues (20). The precise set of transcription factor genes occupied by HNF1 α , HNF4 α , and HNF6, and the extent to which they are co-occupied by the HNF regulators, differed substantially between these two tissues.

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The transcription factor binding data was used to identify regulatory network motifs, simple units of transcriptional regulatory network architecture that suggest mechanistic models (Fig 2B) (4, 23). Our data confirm previous reports that HNF1α and HNF4α occupy one another's promoters in both hepatocytes and islets, forming a multi-component loop (24-26). Multicomponent loops provide the capacity for feedback control and produce bistable systems that can switch between two alternate states (23). It has been suggested that the multicomponent loop present between $HNF1\alpha$ and $HNF4\alpha$ is responsible for stabilization of the terminal phenotype in pancreatic beta cells (26). Applicants also found that HNF6 serves as a master regulator for feedforward motifs in hepatocytes and pancreatic islets involving over 80 genes in each tissue (Figures 20 and 22). For example, in hepatocytes, HNF6 binds the HNF4 α 7 promoter, and HNF6 and HNF4 α together bind *PCK1*, which encodes phosphoenolpyruvate carboxykinase, an enzyme key to gluconeogenesis (Fig 2B). A feedforward loop can act as a switch designed to be sensitive to sustained, rather than transient, inputs (23). HNF1 α , HNF4 α and HNF6 were also found to form multi-input motifs by collectively binding to sets of genes in hepatocytes and islets. This regulatory motif suggests coordination of gene expression through multiple input signals. Applicants also found that HNF6, HNF4α, and HNF1α form a regulator chain motif with THRA (NR1D1); regulator chain motifs represent the simplest circuit logic for ordering transcriptional events in a temporal sequence (4, 23). Additional examples of these regulatory motifs can be found in Figures 20 and 23 (20). Figures 20-24, panels A and B, show transcriptional regulators occupied by HNF transcription factors and their regulatory loops. Figures 4-10 show additional controls and data generated by the experiments described herein.

Our results suggest that the nuclear hormone receptor HNF4 α contributes to regulation of a large fraction of the liver and pancreatic islet transcriptomes by binding directly to almost half of the actively transcribed genes. This likely explains why

- HNF4α is crucial for development and proper function of these tissues (12-15, 17, 18). Perhaps most importantly, our results suggest a mechanistic explanation for the recent discovery that polymorphisms in the islet-specific P2 promoter for the splice variant HNF4α7 can greatly increase the risk of type II diabetes (27-30). Applicants found that multiple HNF factors bind directly to the P2 promoter in primary, healthy human islets.
- Alterations in the binding sites for these factors could cause misregulation of HNF4α expression and thus its downstream targets, leading to beta cell malfunction and diabetes.

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Claims:

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1. A method of determining which genes from a subset of genes are regulated by a transcriptional regulator expressed in a cell, the method comprising

- (a) selectively isolating chromatin from the cell to generate isolated chromatin;
- (b) selectively isolating chromatin fragments from the isolated chromatin to generate bound chromatin fragments, wherein the bound chromatin fragments are bound by the transcriptional regulator;
- (c) amplifying both the bound chromatin fragments to generate amplified chromatin fragments and the isolated chromatin to generate amplified control chromatin;
- (d) hybridizing the amplified control chromatin and the amplified chromatin fragments to a DNA microarray, wherein the DNA microarray comprises
 - (1) at least 10,000 experimental spots, each experimental spot comprising an experimental DNA, each experimental DNA comprising a promoter region from a gene in the subset; and
 - (2) at least 100 control spots, each control spot comprising a control DNA, each control DNA comprising a non-promoter region; and
- (e) determining and comparing a hybridization signal at each of the spots on the microarray between those generated by
 - (1) the amplified control chromatin; and
 - (2) the amplified chromatin fragments;
- wherein a gene in the subset is said to be regulated by the transcriptional regulator in the cell if a spot comprising a promoter region of said gene displays a higher level of hybridization by the amplified chromatin fragments than by the amplified control chromatin.
- The method of claim 1, wherein the level of hybridization of the amplified chromatin fragments to each experimental spot is normalized by the level of hybridization of the amplified chromatin fragments to the control spots.

3. The method of claim 1, wherein the level of hybridization of the amplified chromatin fragments to each experimental spot is normalized by subtracting the mean level of hybridization of the amplified chromatin fragments to the control spots.

- 4. The method of claim 1, wherein the higher level of hybridization comprises at least a two-fold higher level of hybridization.
- The method of claim 1, wherein the transcriptional regulator is native to the cell.
 - 6. The method of claim 1, wherein the transcriptional regulator is not a recombinant transcriptional regulator.

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- 7. The method of claim 1, wherein the cell is a primary cell.
- 8. The method of claim 7, wherein the cell is a human cell.
- 20 9. The method of claim 8, wherein the cell is a transplant-grade human cell.
 - 10. The method of claim 1, wherein step (b) comprises immunoprecipitation of the transcriptional regulator.
- The method of claim 1, wherein step (c) comprises ligation-mediated polymerase chain reaction (LM-PCR).
 - 12. The method of claim 1, wherein the promoter region of the gene comprises from at least 700bp upstream to at least 200 bp downstream of the transcriptional start site of the gene.

13. The method of claim 1, wherein the promoter region comprises at least 30, 40, 50, or 60 or nucleotides in length.

- 14. The method of claim 1, wherein the promoter region of the gene comprises a sequence of at least 30 nucleotides whose sequence is identical to a region stretching from 3 kb upstream to 1 kb downstream of the transcriptional start site of said gene.
- The method of claim 1, wherein the non-promoter region comprises an open reading frame.
 - 16. The method of claim 1, wherein the transcriptional regulator is a basal transcription factor.
- 15 17. The method of claim 16, wherein the transcriptional regulator is an RNA polymerase II or a TATA-binding protein.
- 18. A method of identifying a transcriptional regulatory network in a cell, the method comprising determining if a transcriptional regulator regulates

 20 additional transcriptional regulators in the cell using the method of claim 1, wherein a transcriptional regulatory network is identified if at least one additional transcriptional regulator is determined to be regulated by the transcriptional regulator.
- The method of claim 18, wherein the experimental DNA comprises promoter regions from the additional transcriptional regulators.
 - 20. A method of identifying a transcriptional regulatory network in a cell, the method comprising determining if a transcriptional regulator regulator
- 30 (i) its own promoter; or

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(ii) a promoter from a plurality of transcriptional regulators, using the method of claim 1, wherein the experimental DNA comprises

(a) a promoter from the transcriptional regulator; and

- (b) promoters from the plurality of transcriptional regulators; wherein a transcriptional regulatory network is identified if the transcriptional regulator regulates itself or if it regulates at least one of the plurality of transcriptional regulators.
- 21. A method of identifying transcriptional regulatory networks in a cell, the method comprising

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- (a) determining, by repeating the method of claim 1 for each of a plurality of transcriptional regulators, the genes in a subset which are regulated by each of the plurality of transcriptional regulators, wherein the experimental DNA comprises promoter regions for each of the plurality of transcriptional regulators;
- (b) determining if any one of the plurality of transcriptional regulators are regulated by at least one of the plurality of transcriptional regulators; wherein a transcriptional regulatory network is identified if any one of the plurality of transcriptional regulators is regulated by at least one of the plurality of transcriptional regulators.
- 20 22. The method of claim 21, further comprising determining if a gene is regulated by more than one of the plurality of transcriptional regulators.
 - 23. A DNA microarray for determining promoter occupancy in a human cell, the microarray comprising
 - (1) at least 10,000 experimental spots, each experimental spot comprising an experimental DNA, each experimental DNA comprising a promoter region from a human gene in the subset; and
 - (2) at least 100 control spots, each control spot comprising a control DNA, each control DNA comprising a non-promoter region; wherein at least 75% of the promoter regions comprise from at least 700bp upstream to at least 200 bp downstream of the transcriptional start site.

24. A method of estimating if a transcriptional regulator is a global transcriptional regulator, the method comprising

(a) selectively isolating chromatin from a tissue;

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- (b) identifying promoter regions from the chromatin which are bound by a candidate global transcriptional regulator;
- (c) identifying promoter regions from the chromatin which are bound by a member of the basal transcriptional machinery; and
- (d) comparing the promoter regions identified in steps (b) and (c) to determine the ratio between (i) the number of promoter regions bound by both the candidate global transcriptional regulator and the member of the basal transcriptional machinery; and (ii) the number of promoter regions bound by the member of the basal transcriptional machinery

wherein a transcriptional regulator is a global transcriptional regulator when the ratio is greater than 0.2.

- 25. The method of claim 24, wherein steps (b) and (c) are performed using a DNA microarray.
- 26. The method of claim 25, wherein the DNA microarray comprises
 (i) at least 10,000 experimental spots, each experimental spot comprising an experimental DNA, each experimental DNA comprising a promoter

region from a human gene in the subset; and

- (ii) at least 100 control spots, each control spot comprising a control DNA, each control DNA comprising a non-promoter region;
- 27. The method of claim 24, wherein the member of the basal transcriptional machinery is an RNA polymerase II or a TATA-binding protein.
- 30 28. The method of claim 24, wherein the tissue is transplant-grade tissue.
 - 29. The method of claim 24, wherein the tissue is freshly-isolated human tissue.

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30. The method of claim 29, wherein the tissue is from a subject afflicted with a disorder.

- 5 31. The method of claim 30 wherein the disorder is a hyperplastic condition.
 - 32. A method of identifying at least one target gene for the development of a therapeutic to treat or prevent a disorder in a subject, wherein at least one form of the disorder is caused by an altered activity in a transcriptional regulator or in a suspected transcriptional regulator, the method comprising
 - (a) identifying the genes regulated by the transcriptional regulator in a cell;
 - (b) determining if the transcriptional regulator is a broad-acting transcriptional regulator or a narrow-acting transcriptional regulator, wherein if the transcriptional regulator is a broad acting transcriptional regulator then the transcriptional regulator is a target gene for the development of a therapeutic, and wherein if the transcriptional regulator is a narrow acting transcriptional regulator then
 - (i) determining if at least one gene regulated by the transcriptional regulator is likely causative in the disorder, wherein a gene that is likely causative in the disorder is a target gene for the development of a therapeutic; and
 - (ii) reiterating steps (a) and (b) for at least one gene that is regulated by the transcriptional regulator in the cell and that either
 - (1) encodes a transcriptional regulator or
 - (2) is suspected to encode a transcriptional regulator,with the modification that the transcriptional regulator of steps (a) and(b) is said gene,

thereby identifying at least one target gene for the development of a therapeutic to treat or prevent a disorder in the subject.

33. The method of claim 32, wherein identifying the genes regulated by the

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transcriptional regulator in a cell comprises chromosome-wide location

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- The method of claim 32, wherein identifying the genes regulated by the transcriptional regulator in the cell comprises using the method of claim 1.
 - 35. The method of claim 32, wherein the transcriptional regulator is a master regulatory gene.
- The method of claim 35, wherein the master regulatory gene is SOX1-18, OCT6, PAX3, Myocardin, GATA1-6, TCF1/HNF1A, HNF4A, HNF6, NGN3, C/EBP, FOXA1-3, IPF1, GATA, HNF3, NKX2.1, CDX, FTF/NR5A2, C/EBPbeta, SCL1, SKIN1, or a member of the neurogenin, LK, LMO, SOX, OCT, PAX, GATA or MyoD family of transcription factors.

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analysis.

- The method of claim 32, wherein the transcriptional regulator is PAX3, EGR-1, EGR-2, OCT6, a SOX family member, a GATA family member, a PAX family member, an OCT family member, RFX5, WHN, GATA1, VDR, CRX, CBP, MeCP2, AML1, p53, PLZF, PML, Rb, WT1, NR3C2, GCCR, PPARgamma, SIM1, HNF1alpha, HNF1beta, HNF4alpha, PDX1, MAFA, FOXA2, or NEUROD1.
- 38. The method of claim 32, wherein the cell is derived from a tissue whose function is impaired in the disorder.

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- 39. The method of the claim 32, wherein the broad acting gene regulates at least about 2.5% of the genes in the cell, and wherein the narrow acting gene regulates less than about 2.5% of the genes in the cell.
- The method of claim 32, wherein the gene is suspected to encode a transcriptional regulator if it shares at least 30% amino acid sequence identity with the DNA binding domain of a transcriptional regulator.

41. The method of claim 32, wherein the transcriptional regulator in the cell is a mutant transcriptional regulator.

- 5 42. The method of claim 32, wherein the transcriptional regulator in the cell has altered activity.
- The method of claim 32, wherein the gene regulated by the transcriptional regulator is likely causative of the disorder when a mutation in the gene results in at least one phenotype or symptom associated with the disorder.
 - 44. The method of claim 32, wherein the gene regulated by the transcriptional regulator is likely causative of the disorder when the gene encodes an enzyme or signaling molecule which functions in a pathway that is impaired in the disorder.
 - The method of claim 32, wherein the altered activity in the transcriptional regulator comprises at least one of the following:

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- (a) an alteration in the binding affinity of the transcriptional regulator to DNA;
- (b) an alteration in the ability of the transcriptional regulator to bind to RNA polymerase, to an RNA polymerase holoenzyme, or to a second transcriptional regulator;
- (c) an alteration in the binding affinity of the transcriptional regulator to a ligand;
- (d) an alteration in expression level or expression pattern of the transcriptional regulator; or
- (e) an alteration in an ability of the transcriptional regulator to form homomultimers or heteromultimers.
- 46. The method of claim 32, wherein the disorder is characterized by impaired function of at least one of the following: brain, spinal cord, heart, arteries,

esophagus, stomach, small intestine, large intestine, liver, pancreas, lungs, kidney, urinary tract, ovaries, breasts, uterus, testis, penis, colon, prostate, bone, muscle, cartilage, thyroid gland, adrenal gland, pituitary, bone marrow, blood, thymus, spleen, lymph nodes, skin, eye, ear, nose, teeth or tongue.

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- 47. The method of claim 32, wherein the therapeutic comprises a small molecule drug, an antisense reagent, an antibody, a peptide, a ligand, a fatty acid, a hormone or a metabolite.
- 10 48. The method of claim 32, wherein the subject is a mammal.
 - 49. The method of claim 48, wherein the mammal is a human.
- 50. The method of claim 32, wherein the transcriptional regulator is a transcriptional activator or a transcriptional repressor.
 - 51. The method of claim 32, wherein the transcriptional regulator is native to the cell.
- The method of claim 32, wherein the transcriptional regulator is from a species different from that of the cell.
 - 53. The method of claim 52, wherein the transcriptional regulator is a viral transcriptional regulator.

- A method of treating or preventing type II diabetes in a subject, comprising administering to the subject a therapeutically effective amount of an agent that increases the global transcriptional activity of HNF4alpha.
- 30 55. A method of treating or preventing a disorder associated with low transcriptional activity of HNF4alpha in a subject, comprising administering to the subject a therapeutically effective amount of an agent that increases the

global transcriptional activity of HNF4alpha.

56. A method of treating or preventing a disorder associated with high transcriptional activity of HNF4alpha in a subject, comprising administering to the subject a therapeutically effective amount of an agent that decreases the global transcriptional activity of HNF4alpha.

- 57. A method of increasing the global transcriptional activity in a liver or a pancreatic cell comprising contacting the cell with an agent which increases the global transcriptional activity of HNF4alpha.
- A method of decreasing the global transcriptional activity in a liver or a pancreatic cell comprising contacting the cell with an agent which decreases the global transcriptional activity of HNF4alpha.

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- A method of regulating the expression level of any one of the genes in Figure 13 in a hepatocyte, the method comprising contacting the cell with an agent which regulates the transcriptional activity of HNF1alpha.
- A method of regulating the expression level of any one of the genes in Figure 14 in a pancreatic cell, the method comprising contacting the cell with an agent which regulates the transcriptional activity of HNF1alpha.
- A method of regulating the expression level of any one of the genes in Figure 16 in a hepatocyte, the method comprising contacting the cell with an agent which regulates the transcriptional activity of HNF6.
 - A method of regulating the expression level of any one of the genes in Figure 17 in a pancreatic cell, the method comprising contacting the cell with an agent which regulates the transcriptional activity of HNF6.
 - 63. A method of regulating the expression level of any one of the genes in Figure

18 in a hepatocyte, the method comprising contacting the cell with an agent which regulates the transcriptional activity of HNF4alpha.

- A method of regulating the expression level of any one of the genes in Figure 19 in a pancreatic cell, the method comprising contacting the cell with an agent which regulated the transcriptional activity of HNF4alpha.
 - A method of identifying transcriptionally active genes that are regulated by a transcriptional regulator in a cell, the method comprising
 - (a) selectively isolating chromatin from a tissue;
 - (b) identifying promoter regions from the chromatin that are bound by the transcriptional regulator;
 - (c) identifying promoter regions from the chromatin that are bound by a member of the basal transcriptional machinery; and
 - (d) comparing the promoter regions identified in steps (b) and (c) to determine overlapping genes,

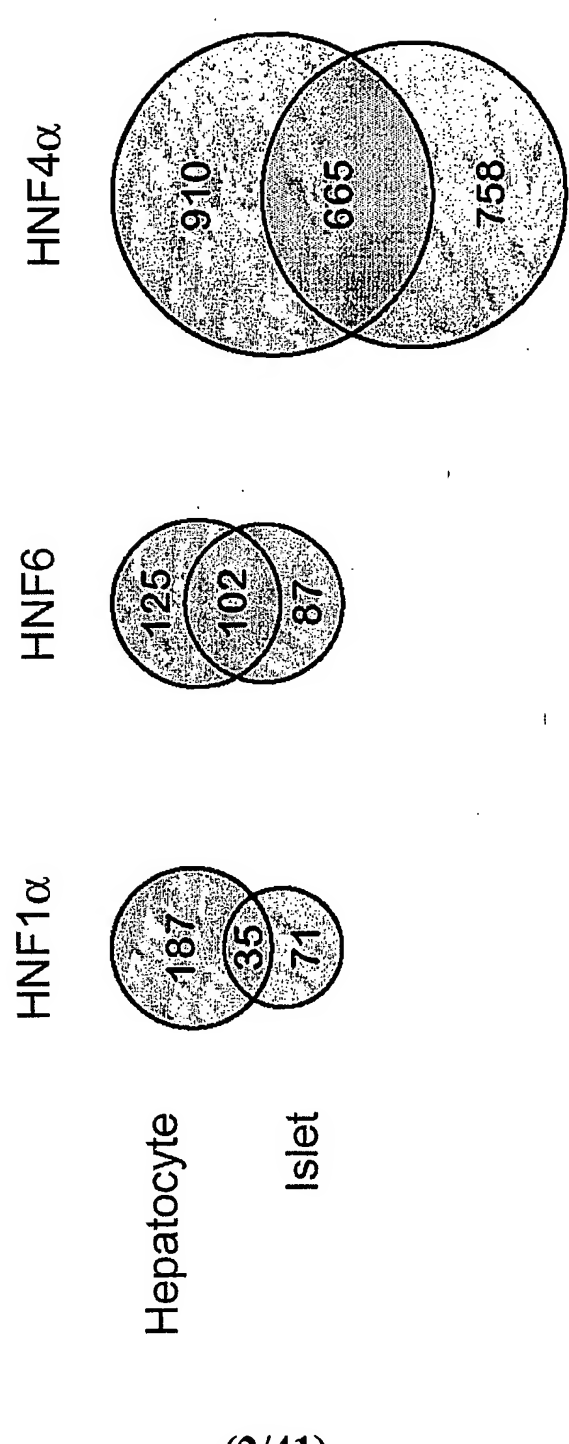
wherein the overlapping genes are transcriptionally active genes regulated by the transcriptional regulator.

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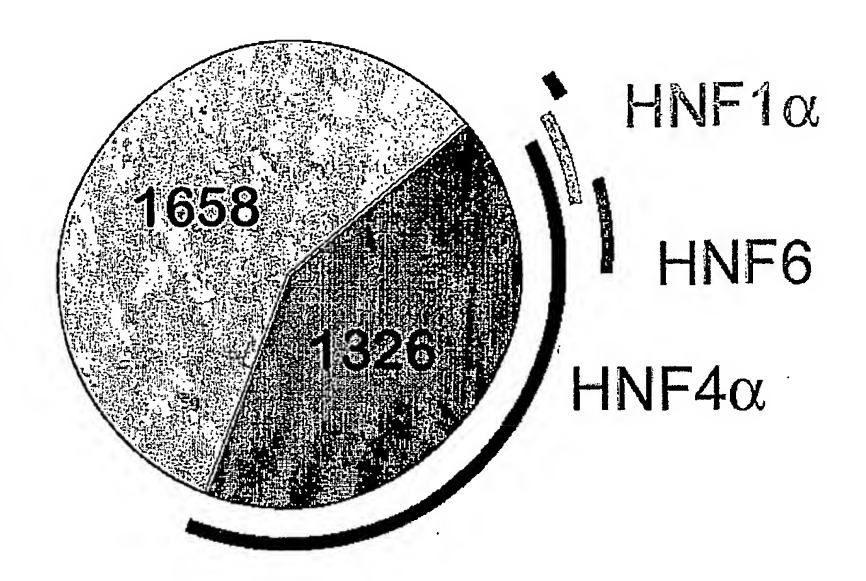
Fig. 1A Hybridize to promoter microarray Chromatin IP to enrich promoters bound by regulator in vivo Purify single cell type liver or pancreas Primary fissue, (1/41)

Fig. 1B

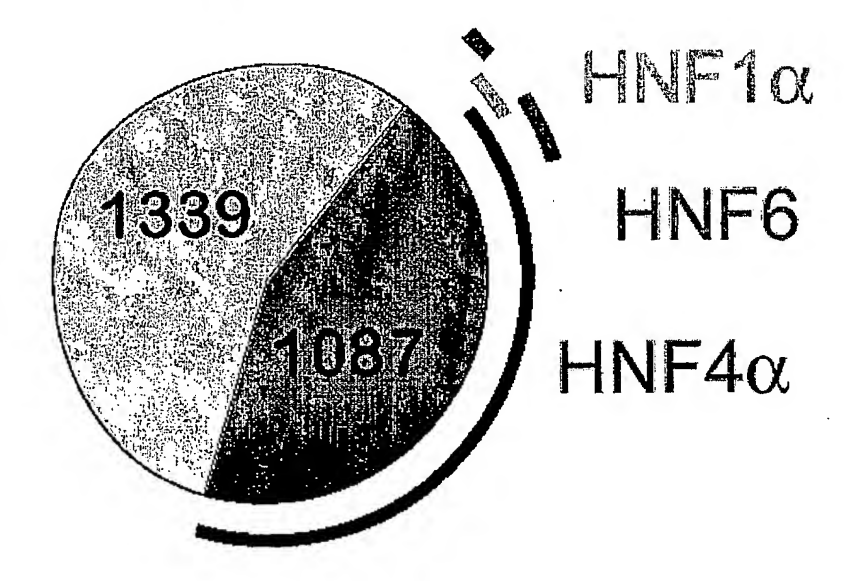


(2/41)

Fig. 1C



Hepatocyte

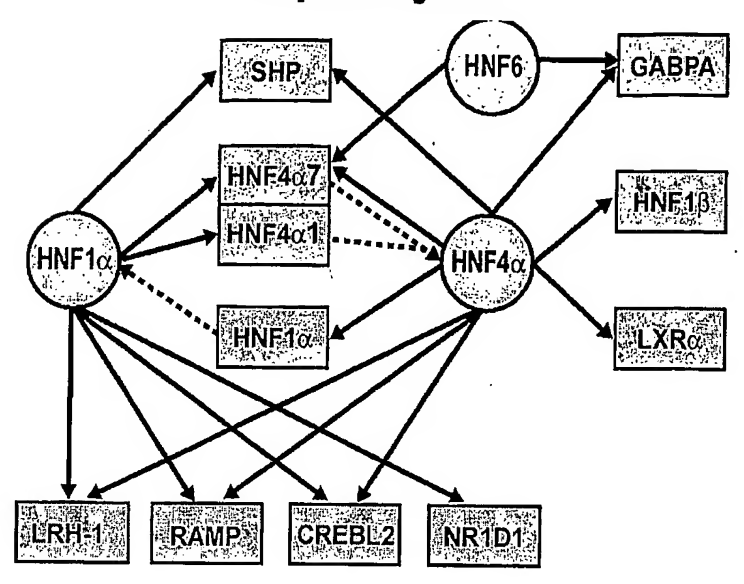


Pancreatic Islet

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Fig. 2A

Hepatocytes



Pancreatic Islets

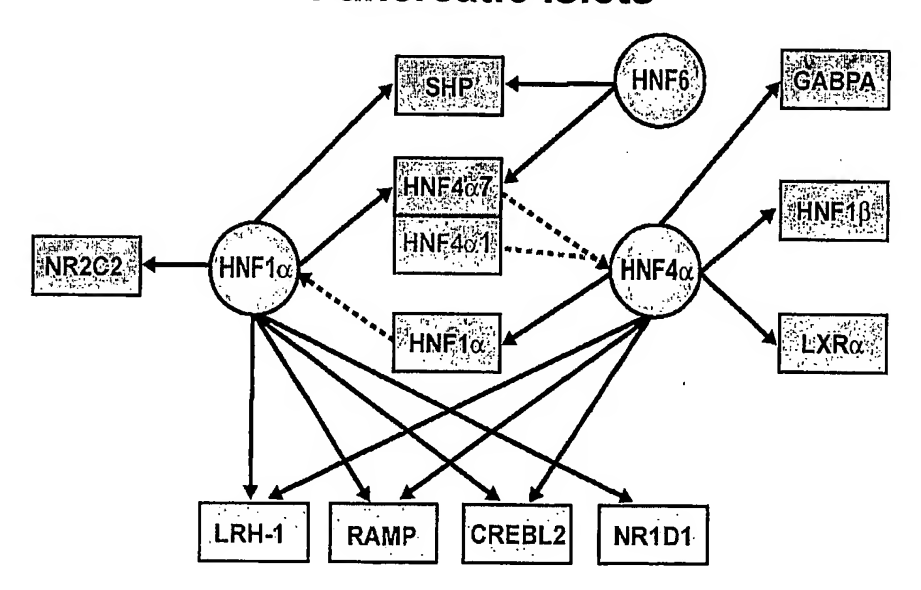
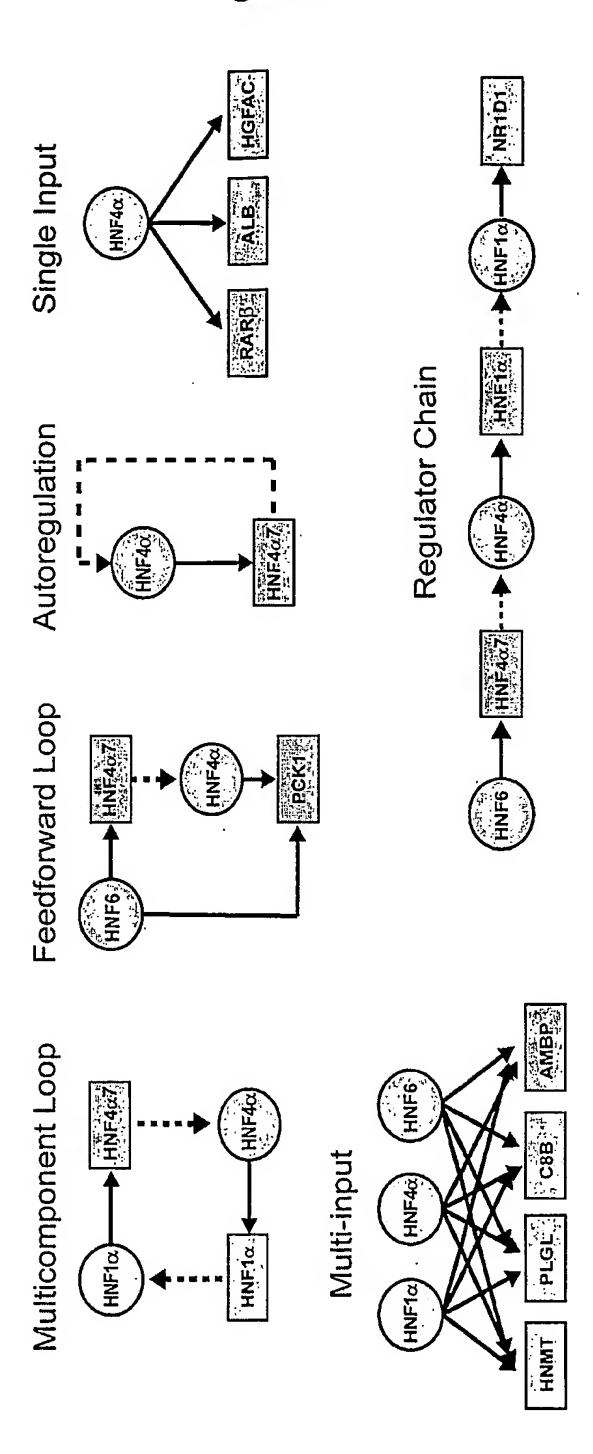


Fig. 2B



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Fig. 3

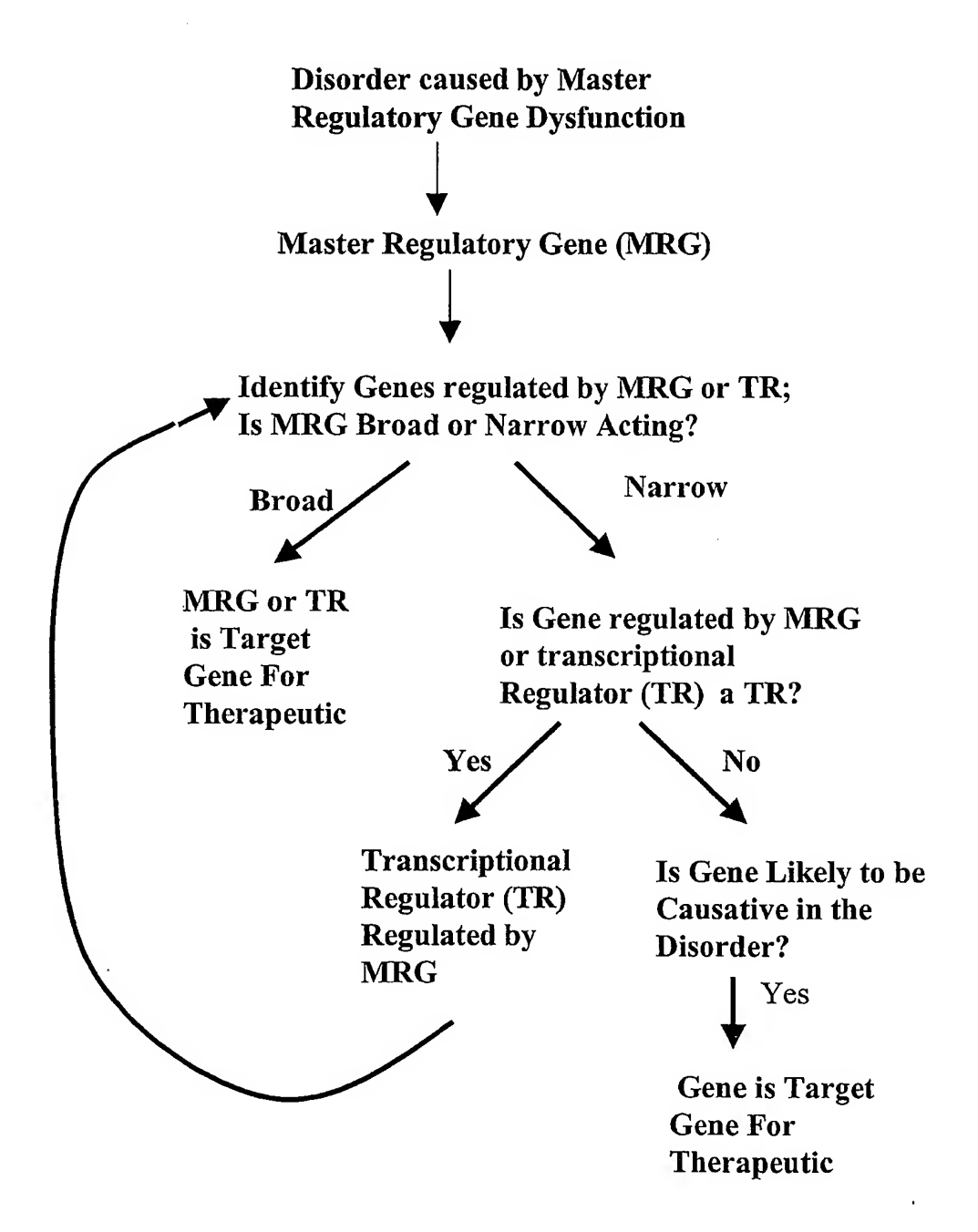


Fig. 4

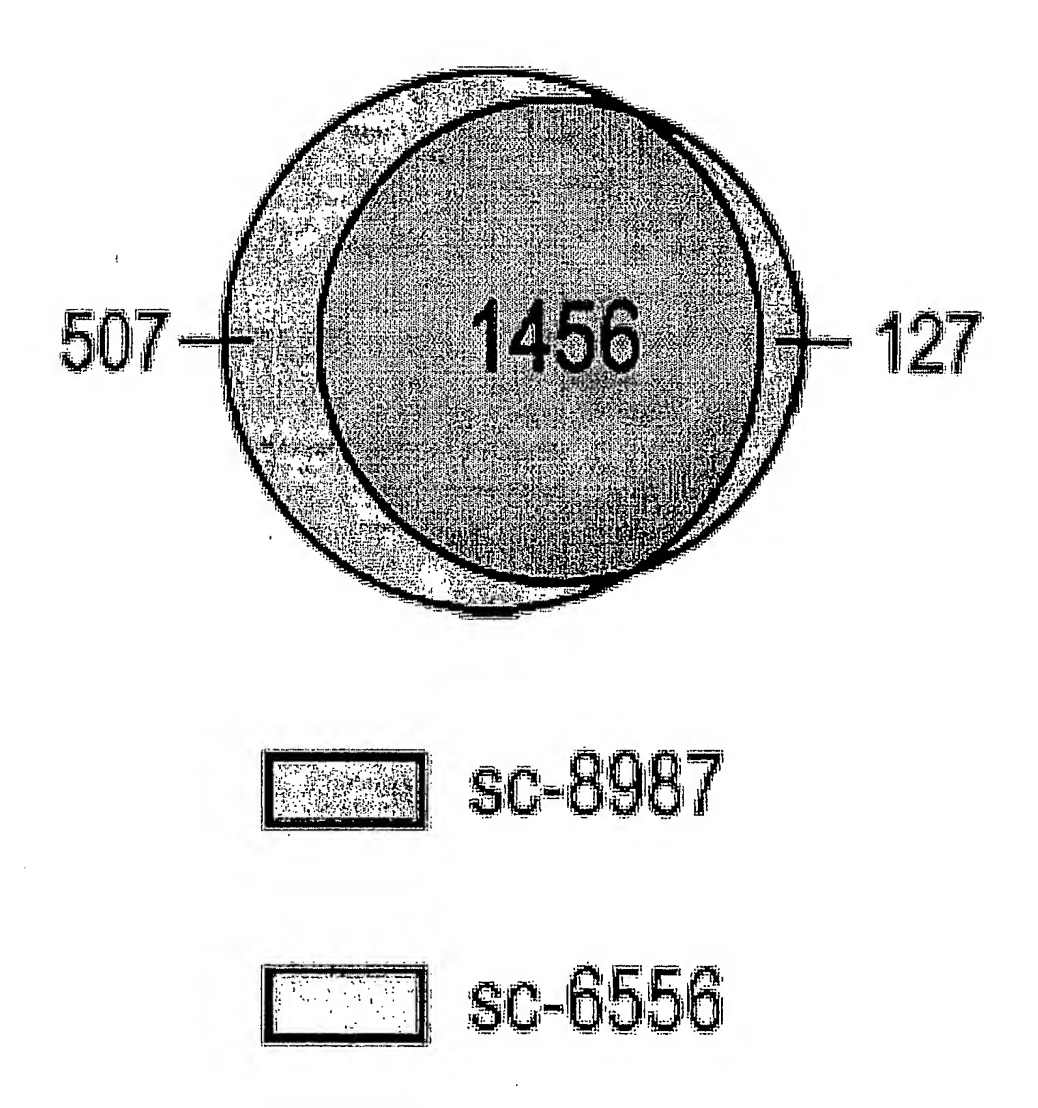
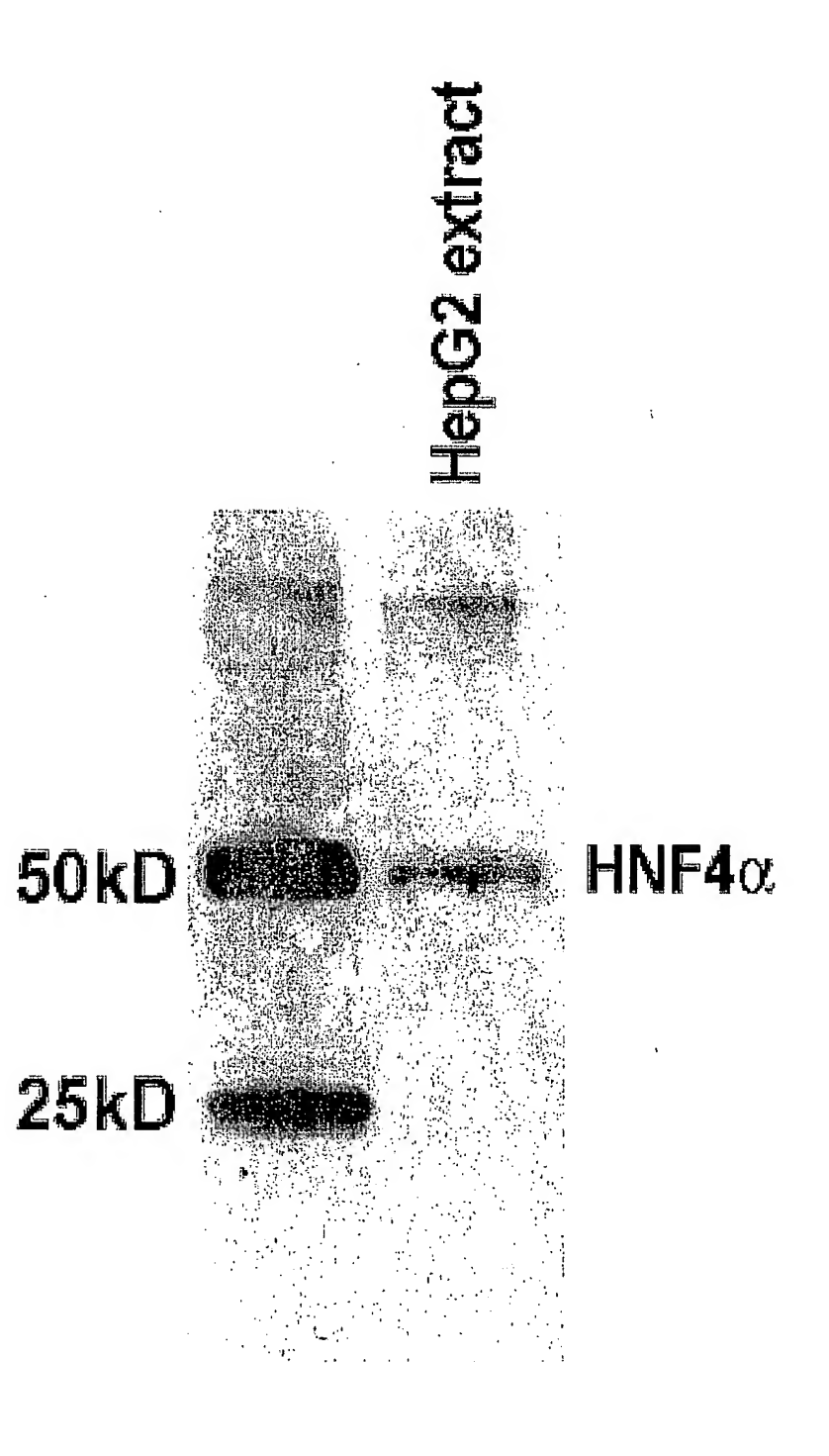


Fig. 5



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Fig. 6A

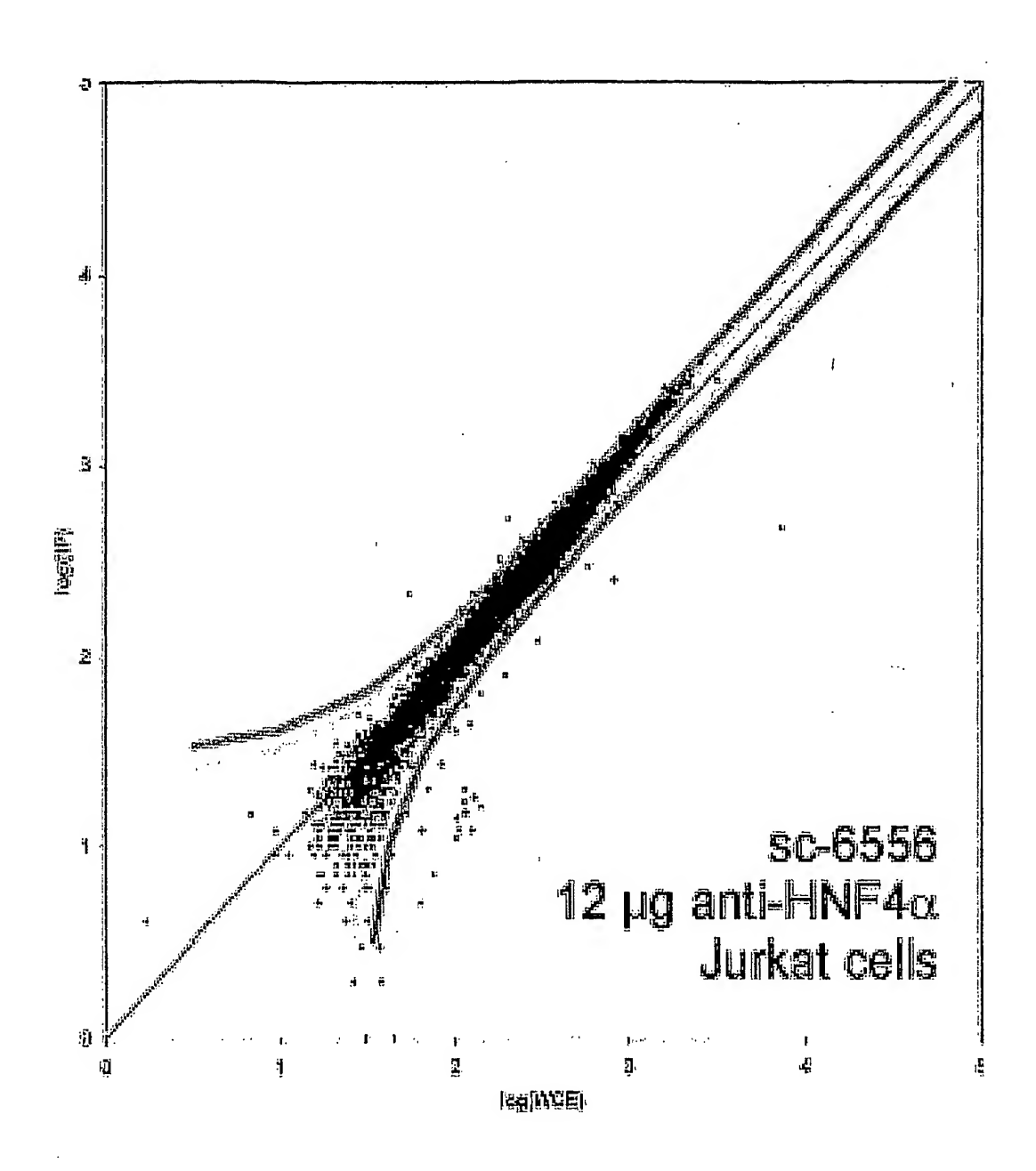


Fig. 6B

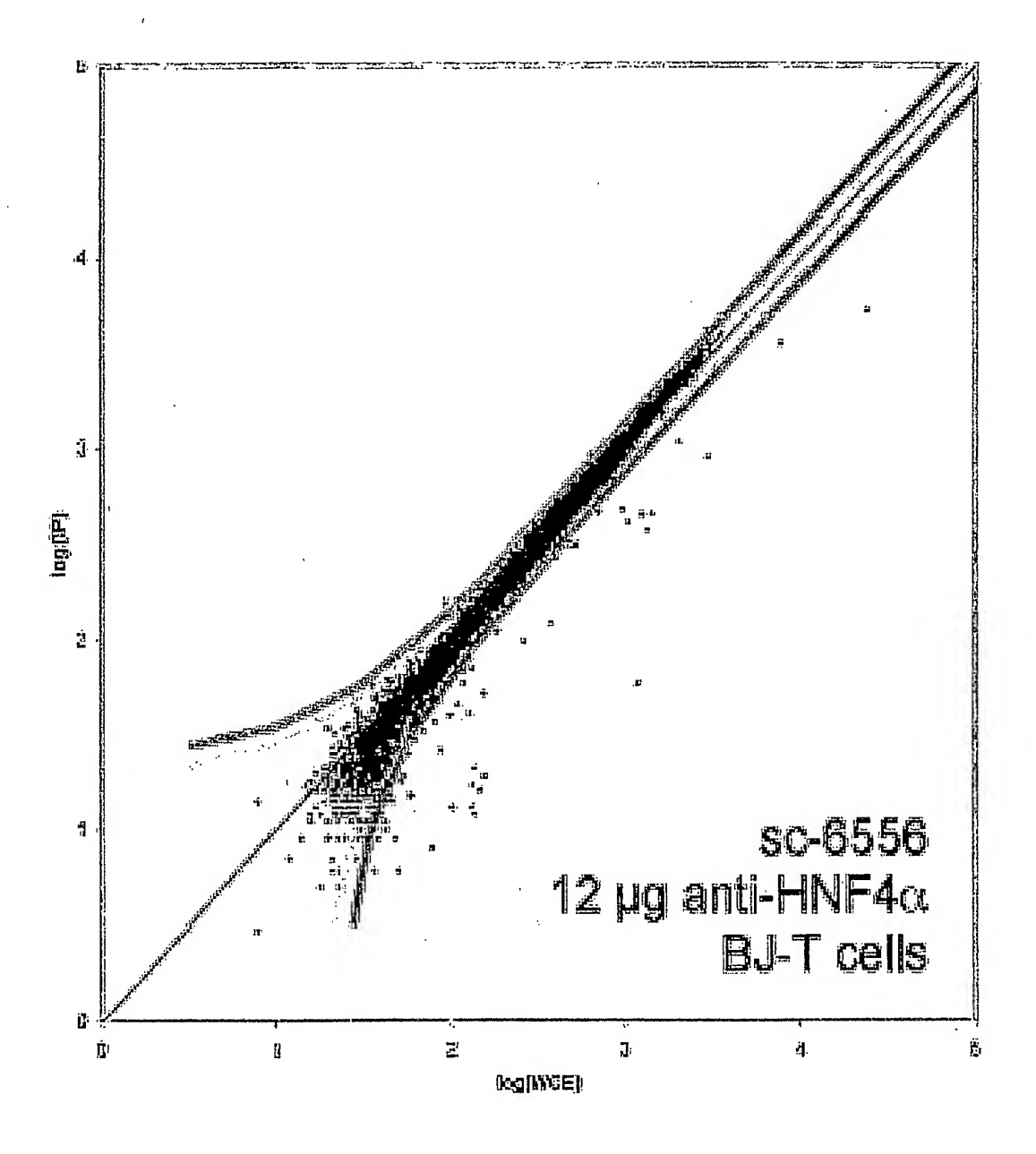


Fig. 6C

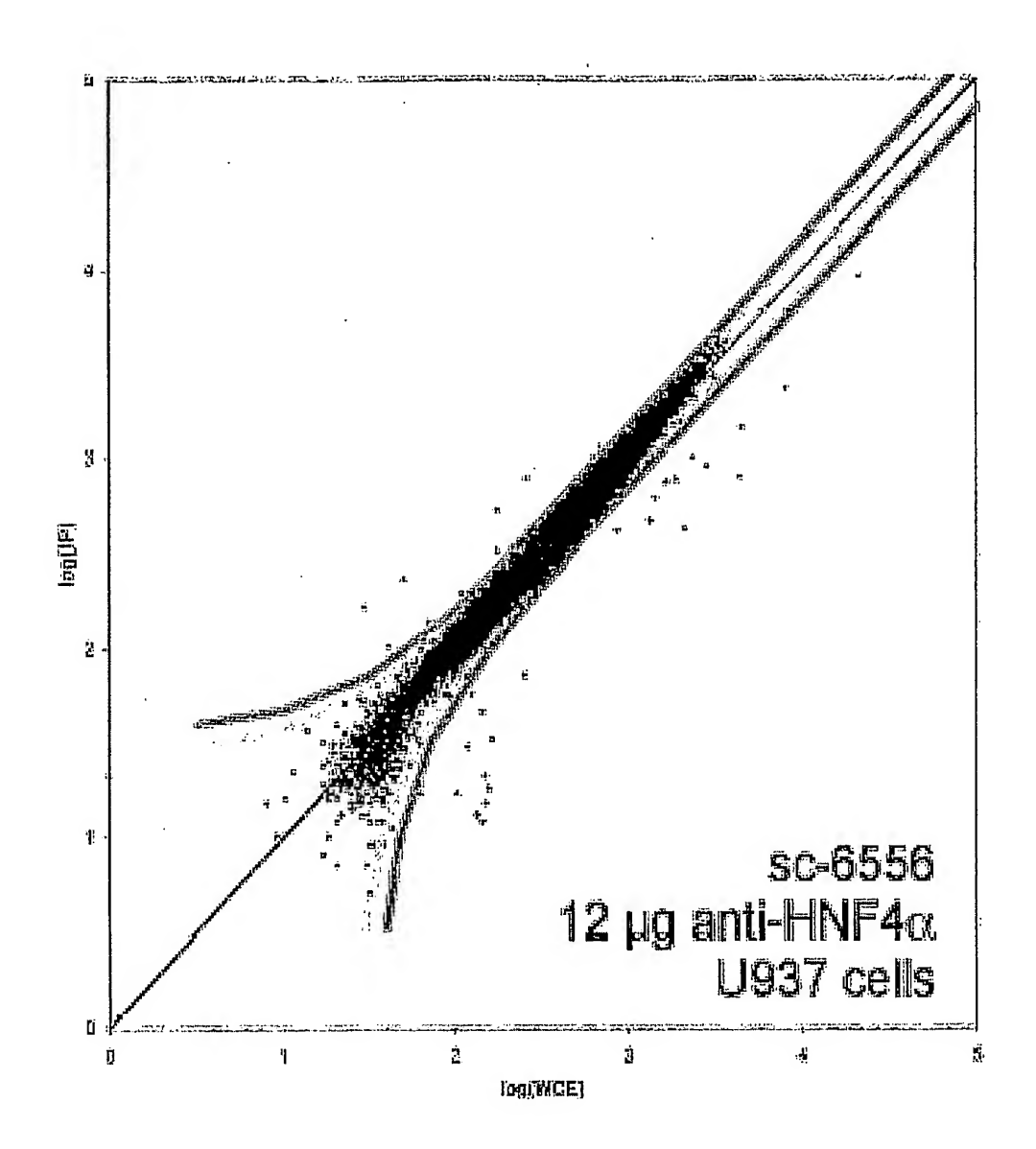
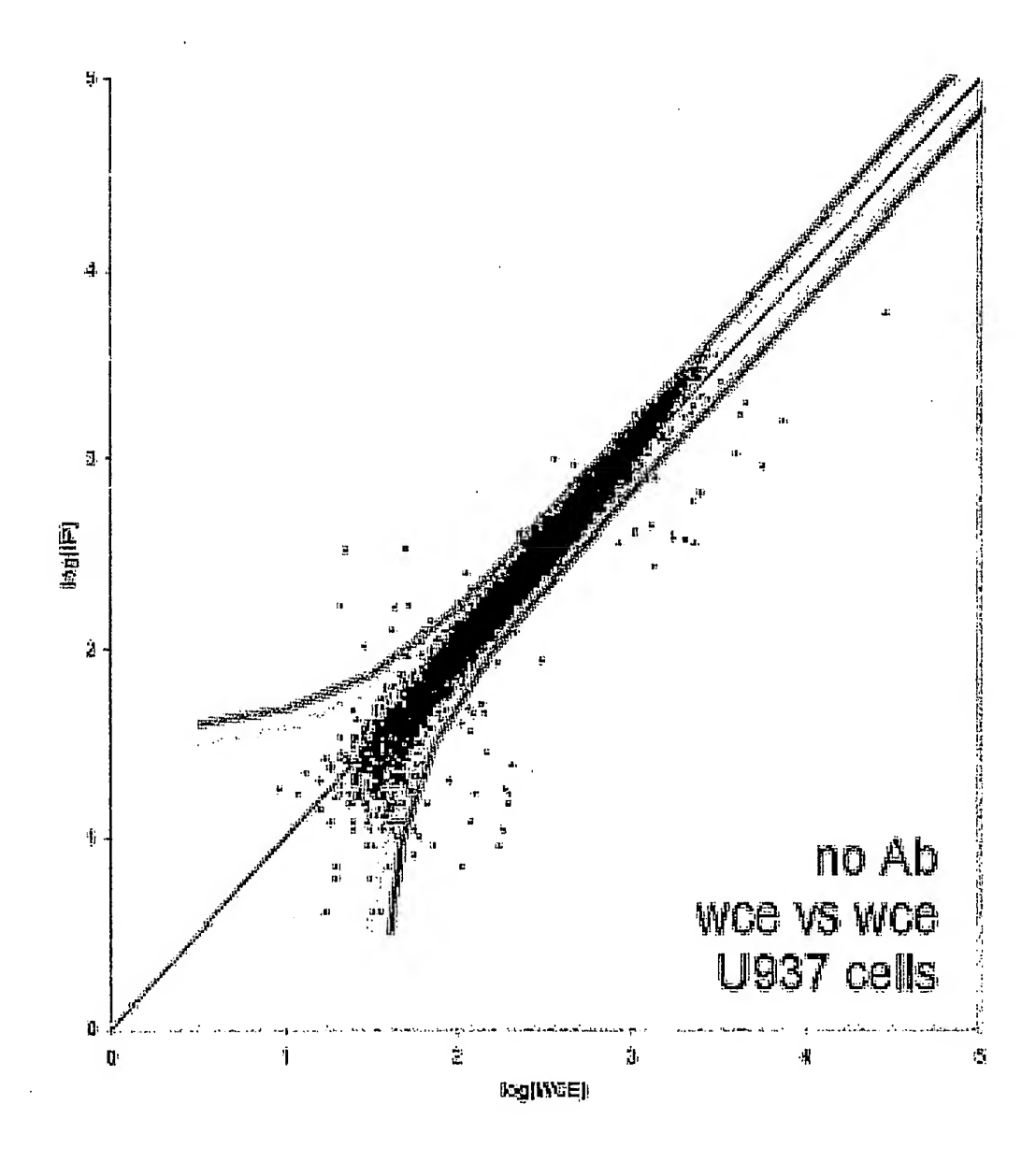
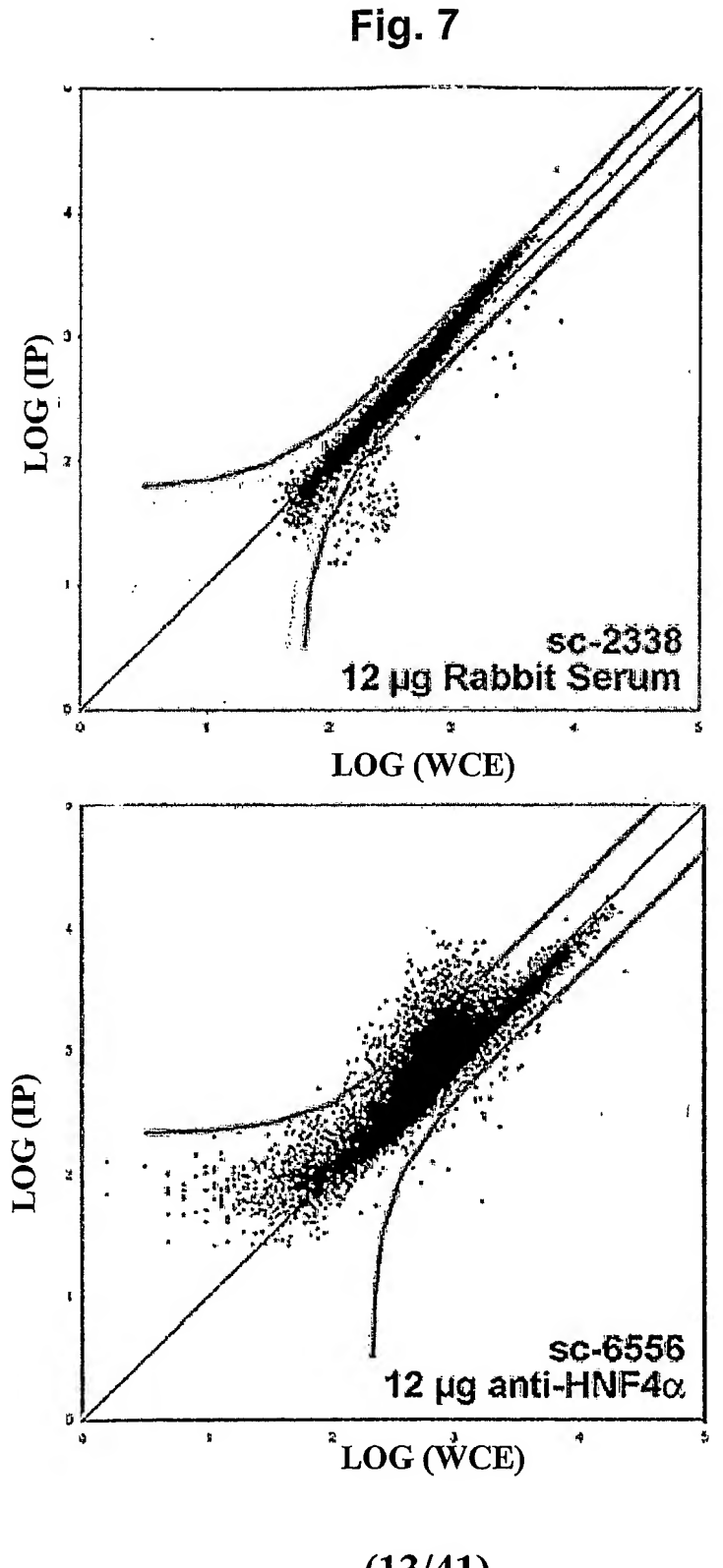


Fig. 6D



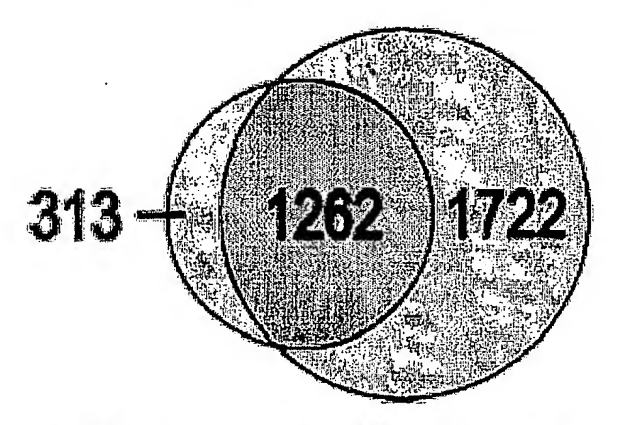
(12/41)



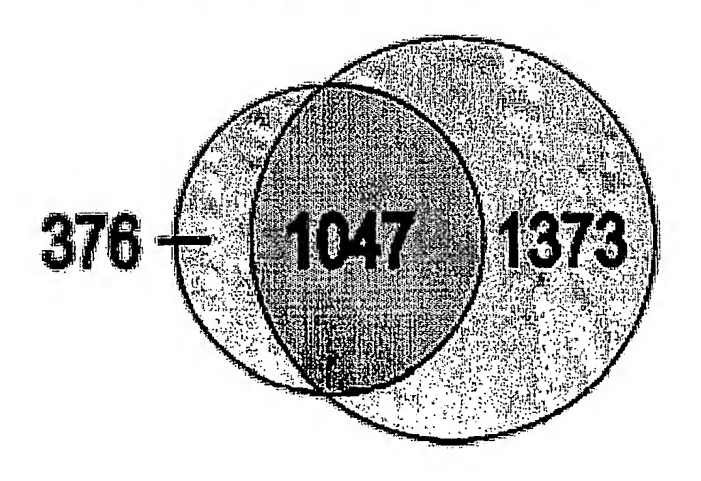
(13/41)

Fig. 8

Hepatocytes



Pancreatic Islets

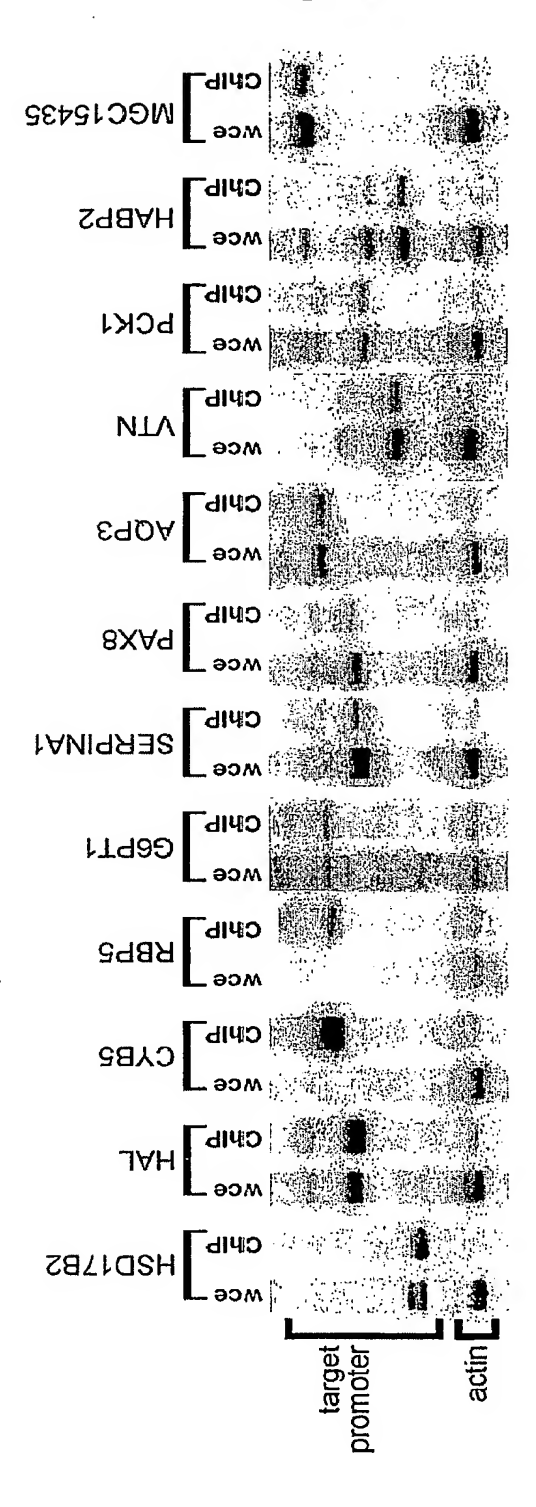


 \square HNF4 α

☐ RNA Pol II

(14/41)

Fig. 9



(15/41)

Fig. 10

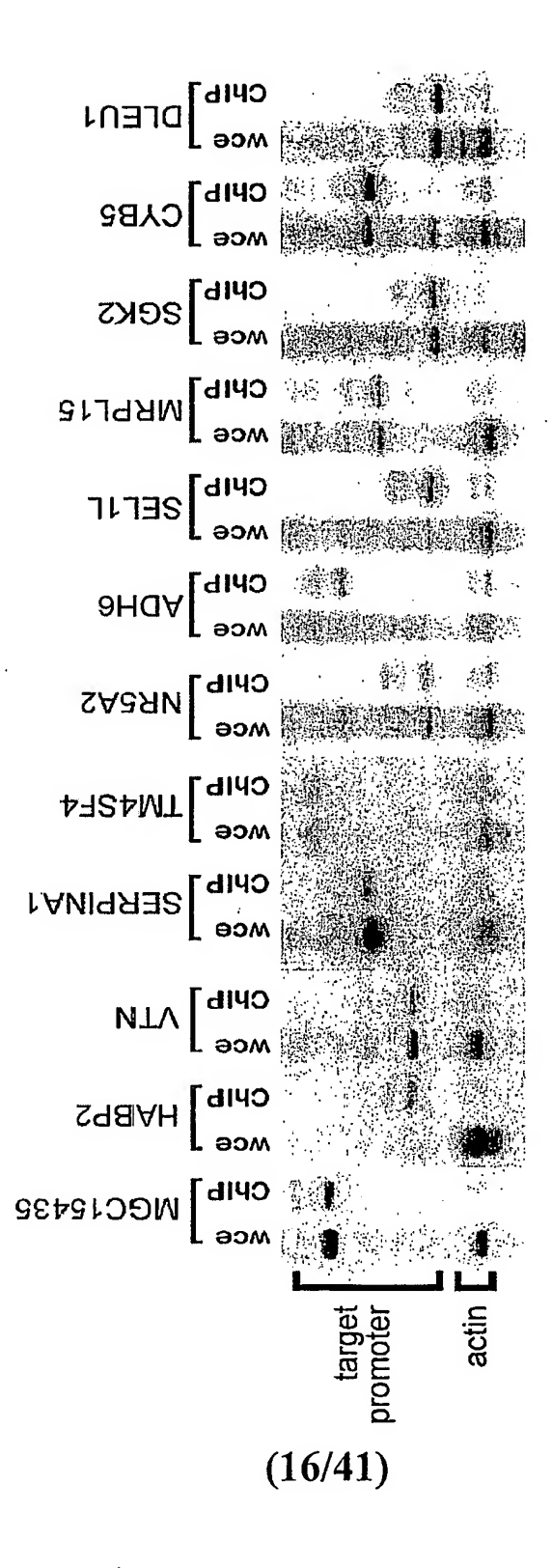


Fig. 11

			Φ					0)	
			ocyt					Scyle	
Name	RefSeq	Description	Hepatocyte	Islets	Name	RefSeq	Description	Hepatocyte	Islets
Chaperone					Signal Trans	duction-Other			
C4BPA	NM_000715	complement 4 binding protein a	¥	4	BIKE	NM_017593	BMP-2 inducible kinase		v
APCS	NM_001639	amyloid P component	•		SGK2	NM_016276	serum/glucocorticold reg. kinase 2	•	•
F11	NM_019559	coagulation factor XI	• 🗸		SEL1L	NM_005065	suppressor of lin-12-like	•	~
C1S	NM_001734	complement component 1s	Y		SCYE1	NM_004757	small cytokine E1	•	
VTN	NM_000638	somatomedin B	V		ANGPTL3	NM_014495	angiopoietin-like 3	V	
EnzymeHyd						ductionRecepto			
PGCP	NM_016134	glutamate carboxypeptidase			HAVCR-1	NM_012206	hepatitis A virus cellular receptor 1		J
gla Lipa	NM_000169 NM_000235	galactosidase, alpha lipase A		v	TACR3 GNB2L1	NM_001059	tachykinin receptor 3		
SPO11	NM_012444	SPO11-like		J	INSR	NM_006098 NM_000208	GTP binding protein, beta2L1		•
PAFAH2	NM_000437	platelet-activating factor 2	v	V	SSTR1	NM_001049	insulin receptor somatostatin receptor 1		y
AADAC	NM_001086	arylacetamide deacetylase		,	TM4SF4	NM_004617	transmembrane 4-4		J
PS-PLA1	NM_015900	phospholipase A1alpha		.	ASGR2	NM_001181	asialoglycoprotein receptor 2	J	•
VNN3	NM_018399	vanin 3	•	v	GPR39	NM_001508	G protein-coupled receptor 39	Ĵ	
CPB2	NM_016413	carboxypeptidase B2	•		IFNAR1	NM_000629	interferon receptor 1	v	
ANPEP	NM_001150	alanyl aminopeptidase	•		TFRC	NM_003234	transferrin receptor	•	
HGFAC	NM_001528	HGF activator	¥		Transcription				
ENPEP	NM_001977	glutamyl aminopeptidase	•		ZNF300	NM_052860	kruppel-like zinc finger protein		¥
Enzyme-Liga					BCL6	NM_001706	B-cell CLL/lymphoma 6		J
MCCC1	NM_020166	methylcrotonoyl-CoA carboxylase		✓	ZNF155	NM_003445	zinc finger protein 155		•
GARS	NM_002047	glycyl-tRNA synthetase	•		FBXO8	NM_012180	F-box only protein 8		¥
TARS	NM_003191	threonyl-tRNA synthetase	•		NR0B2	NM_021969	Small heterodimer protein	~	•
Enzyme-Lya					HNF4a7	AF509467	HNF4alpha, alternate splice	¥	J
UROD	NM_000374	uroporphyrinogen decarboxylase		•	NR5A2	NM_003822	LRH-1/FTZ-F1	•	~
PCK1	NM_002591	PEPCK1	V		ELF3	NM_004433	E74-like factor 3	•	~
HPCL2 HAL	NM_012260 NM_002108	2-hydroxyphytanoyl-CoA lyase histidine ammonia-lyase	.45		NR1D1	NM_021724	THRA1		
FH	NM_000143	fumarate hydratase	v		ATF2 · CREBL2	NM_001880 NM_001310	activating transcription factor 2 CREB-like 2		
EnzymeOxid	_	idinorate frydratase	•		RARB	NM_016152	RAR-beta	Š	
COQ7	NM_016138	COQ7 coenzyme Q, 7		J.		Channel/Pore	1017-900	•	
ADH4	NM_000670	alcohol dehydrogenase 4		Ú	SLC17A2	NM_005835	vesicular glutamate transporter	y	
UQCRC2	NM_003366	ubiqcyt. c reductase core prot. Il	•	•	AQP3	NM_004925	aquaporin 3	J	
CYB5-M	NM_030579	cytochrome b5	•	✓	SLC22A11	NM_018484	hOAT4	•	
CYP2E	NM_000773	cytochrome P450, IIE	4		GJB1	NM_000166	gap junction protein, beta 1	•	
CYB5	NM_001914	cylochrome b-5	✓		Transporter	Lipids and Small	.		
HSD17B2	NM_002153	hydroxysterold dehydrogenase 2	✓		APOH	NM_000042	apolipoprotein H	•	•
ADH1A	NM_000667	alcohol dehydrogenase 1A	•		ALB	NM_000477	albumin	•	
EnzymeTran				•	ABCC2	NM_000392	canalicular OAT	•	
GCNT3	NM_004751	glucosaminyl transferase 3		•	G6PT1	NM_001467	glucose-6-phosphatase, transport	¥	
FNTB	NM_002028	farnesyltransferase beta	.	y	Transporter		mamo: A P At		
HNMT	NM_006895	histamine N-methyltransferase	~		RAB6KIFL	NM_005733	RAB6 interacting, kinesin-like		~
GOT1 UGT2B15	NM_002079 NM_001076	aspartate aminotransferase 1	.4		PEX13	NM_002618	peroxisome biogenesis factor 13		*
GBE1	NM_000158	UDP glycosyltransferase 2B15 glycogen branching enzyme	J		TMP21 RAB33B	NM_006827 NM_031296	transmembrane trafficking protein RAS oncogene	.4	V
Enzyme Regu		grycogen branching enzyme	·		NAPA	NM_003827	alpha SNAP		•
SERPING1	NM_000062	C1-Inhibitor	v		AP3M1	NM_012095	adaptor-related prot. Complex	Ĵ	
SERPINAT	NM_000295	alpha-1-antitrypsin	•		SNX17	NM_014748	sorting nexin 17	•	
ITIH4	NM_002218	inter-alpha inhibitor H4	✓		_,,,,,,				
AHSG	NM_001622	alpha-2-HS-glycoprotein	✓	•					
Ligand Bindlr									
TMOD2	NM_014548	tropomodulin 2		•					
IGF8P1	NM_000596	IGF binding protein 1	•						
MT1X	NM_005952	metallothionein 1X	~						
CRP	NM_000567	C-reactive protein	•						
APOA2	NM_001643	apolipoprotein A-II	✓						

Fig. 12

	BJ-T vs	BJ-T vs Hepatocytes*	BJ-T vs Pancreatic Islets*	reatic Islets*
	BJ-T specific genes	Hepatocyte specific genes	BJ-T specific genes	Islet specific genes
HNF4a/RNA Pol II	19/492 (4%)	996/2389 (42%)	29/546 (5%)	825/1898 (43%)
HNF1a/RNA Pol II	2/492 (.4%)	123/2389 (5.1%)	4/546 (.9%)	32/1898 (1.7%)
HNF6/RNA Pol II	7/492 (1.4%)	105/2389 (4.4%)	3/546 (.5%)	68/1898 (3.6%)

Fig. 13

	RefSeq	~							RefSeq
AADAC	NM_001086	DLEU1	NM_005887	HPX	NM_000613	PHF2	NM_005392	ZNF288	NM_015642
ABCC2	NM_000392	DUSP6	NM_022652	HSD11B1	NM_005525	PIST	NM_020399	ZNF361	NM_018555
ACF	NM_014576	EIF4EBP2	NM_004096	HSD17B2	NM_002153	PLCB1	NM_015192		
ADH1A	NM_000667	ELF3	NM_004433	HSPC111	NM_016391	PLG	NM_000301		
ADH1B	NM_000668	ENPEP	NM_001977	HSPC129	NM_016396	PLGL	NM_002665		
ADH6	NM_000672	F11	NM_019559	IFNAR1	NM_000629	PS-PLA1	NM_015900		•
AGT	NM_000029	FE65L2	NM_006051	IGF1R	NM_000875	PZP	NM_002864		
AHSG	NM_001622	FH	NM_000143	IGFBP1	NM_000596	RAB33B	NM_031296	ł	
AK2	NM_001625	FKSG87	NM_032029	INADL	NM_005799	RAMP	NM_016448		
AKR1C2	_	FLJ10242	NM_018036	ITIH3	NM_002217	RARB	NM_016152		
AKR1C3	_	FLJ10276	NM_018045	ITIH4	NM_002218	RBP5	NM_031491		
AKR1C4	-	FLJ10525	NM_018126	ITM2B	NM_021999	RNGTT	NM_003800		
ALB	***	FLJ10583	_	KIAA0022	NM_014880	RPL37AP1	NG_000988	Ì	
ALDH3A2		FLJ10650		KIAA0669	NM_014779	SAC	NM_018417	}	
ALS2	NM_020919	FLJ10774	NM_024662	KIAA0844	NM_014951	SCYE1	NM_004757		
AMBP	NM_001633	FLJ11000		KIAA0872	NM_014940	SEL1L	NM_005065		
ANGPTL3		FLJ11838	=	ľ	-	SERPINA1			
ANPEP	NM_014495	FLJ12788	_	KIAA1041	NM_014947		NM_000295		
AP3M1	NM_001150	FLJ13448	NM_022492	KNG LBP	NM_000893	SERPINA 10	NM_016186		
	NM_012095		_	F .	NM_004139	SERPINA6	NM_001756		
APCS	NM_001639	FLJ13611			NM_015913	SERPINC1	NM_000488		
APG3		FLJ14356	- :	1	NM_016001	SERPINE1	NM_000602		
APOA2	NM_001643	FLJ20080			NM_016632	SERPING1	NM_000062		
APOH	NM_000042	FLJ20718			NM_019043	SGK2	NM_016276		
AQP3		FLJ21272			NM_020143	SLC17A2	NM_005835		
AQP9	_	FLJ21934	NM_024743		NM_021211	SLC22A11	NM_018484		
ARHGAP11A				LY6E	NM_002346	SLPI	NM_003064		
ASGR1	_	FLJ23259	_	M17S2	NM_031858	SNX17	NM_014748		
ASGR2	NM_001181	FNTB	_	M96	NM_007358	SRI	NM_003130		
ATF2	NM_001880	G0S2			_	SSA2	NM_004600		
AUTL1	NM_032852	G3A			NM_031477	SSTR1	NM_001049		
BAT3	NM_004639	G6PT1			NM_031453	SSTR4	NM_001052	İ	
BIKE	NM_017593	GARS	== -			STRAIT11499	NM_021242		
BTN2A1	NM_078476	GBE1		h e	NM_032687	SUPV3L1	NM_003171		
C1S	NM_001734	GCKR	NM_001486	MGC15435	NM_032367	SYN3	NM_133632		
C2	NM_000063	GDI2	NM_001494	MGC955	NM_024097	TARS	NM_003191		
C4BPA	NM_000715	GIOT-2	NM_016264	MIA2	NM_054024	TBPL1	NM_004865	•	
C8B	NM_000066	GJB1	NM_000166	MRPL15	NM_014175	TEF	NM_003216		
CCNE1	NM_001238	GOT1	NM_002079	MRPS18B	NM_014046	TFRC	NM_003234		
CDCA1	NM_031423	GPR39	NM_001508	MSH6	NM_000179	TIEG2	NM_003597		
CISH	NM_013324	GPX2		MT1H		TIEG2	NM_003597		
CLYBL	NM_138280	GRHPR		MT1L	NM_002450	TM4SF4	NM_004617	ľ	
CNTNAP2	NM_014141	GTF2B	_	MT1X	- -	TMEM1	NM_003274		
CPB2		GTF2E1	-	MTHFD1	. —	TNFRSF6	NM_000043		
CREBL2	NM_001310	GTPBG3	-	MTP	NM_000253	UGT1A1	NM_000463		
CRP	NM_000567	HABP2	· · · ·	NAPA	_	UGT2B11	NM_001073		
CTSZ	NM_001336	HAL	-	NET-2	NM_012338	UGT2B15	NM_001076		
CYB5	NM_001914	HAO1		NFKBIB	NM_002503	UQCRC2	NM_003366		
CYB5-M	NM_030579	HCAP-G	-	NPC1L1	NM_013389	VNN3	NM_018399		
CYP2E	NM_000773	HGD	_	NR082	NM_021969	VTN	NM_000638	,	
CYP3A43	NM_022820	HGFAC		NR1D1	NM_021724	WBP4	NM_007187		
DAF	NM_000574	HNF4A		NR5A2	_	WDF2	NM_052950		
DC13		HNF4A		NRD1	-	WDF2 WDR12			
	-		-		_		NM_018256		
DKFZP564O0463 DKFZP586A0522		HNF4a7 HNMT	AF509467 NM_006895	PAFAH2 PAX8	NM_000437	XDH XPC	NM_000379		
					NM_013952		NM_004628		
DKFZP586M0122	NM_U15425	HPCL2	NM_012260	PCK1	NM_002591	ZK1	NM_005815	}	

Fig. 14

Name	RefSed	Name Name	RefSeq 35 4 3 34
AADAC	NM_001086	KIAA0101	NM_014736
ABCC9	NM_020297	KIAA0399	NM_015113
ADH4	NM_000670	KIAA0844	NM_014951
APOH	NM_000042	KIF13A	NM_022113
ARHGAP11A	NM 014783	KIR-023GB	NM_015868
B29	NM_031939	KIR2DS2	NM_012312
BCL6	NM_001706	KIR3DL1	NM_013289
BIKE	NM_017593	KRTAP1.1	NM_030967
C4BPA	NM_000715	KRTHA3A	NM_004138
C6orf11	NM_005452	LIPA	NM_000235
CDC45L	NM_003504	LOC113201	NM_138423
COL3A1	NM_000090	LOC113220	NM_138424
COQ7	NM 016138	LOC51092	
CPXCR1		LOC51092	NM_015996
	NM_033048	3	NM_020147
CRH	NM_000756	MCCC1	NM_020166
CTSZ	NM_001336	MGC10500	NM_031477
CYB5-M	NM_030579	MGC15677	NM_032878
DKFZP564J157	NM_018457	MIA2	NM_054024
DLEU1	NM_005887	MRPL15	NM_014175
DOCK1	NM_001380	Nod1(-)6kb	NM_006092
DSC1	NM_024421	NPY2R	NM_000910
EIF3S6	NM_001568	NR0B2	NM_021969
ELF3	NM_004433	NR2C2	NM_003298
FBXO8	NM_012180	NR5A2	NM_003822
FE65L2	NM_006051	PAFAH2	NM_000437
FIL1(EPSILON)	NM_014440	PAX8	NM_013952
FLJ10242	NM_018036	pcnp	NM_020357
FLJ10252	NM_018040	PEX13	NM_002618
FLJ10474	NM_018104	PGCP	NM_016134
FLJ10650	NM_018168	PRO2032	NM_018615
FLJ11301	NM_018385	PSMA5	NM_002790
FLJ13273	NM_024751	PS-PLA1	NM_015900
FLJ13385	NM_024853	RAB33B	NM_031296
FLJ13448	NM_025147	RAB6KIFL	NM_005733
FLJ14855	NM_033210	SDCCAG10	NM_005869
FLJ20156	NM_017691	SEL1L	NM_005065
FLJ20225	NM_019062	SGK2	NM_016276
FLJ20234	NM_017720	SLC26A7	NM_052832
FLJ20298	NM_017752	SPO11	NM_012444
FLJ20643	NM_017916	SRI	NM_003130
FLJ20731	NM_017946	SSTR1	NM_001049
FLJ21272	NM 025032	TACR3	NM_001059
FLJ22559	NM_024928	TM4SF4	NM_004617
FNTB	NM_002028	TMOD2	NM_014548
GCNT3	NM_004751	TMP21	NM_006827
GIOT-2	NM_016264	UQCRC2	NM_003366
GLA	NM_010204 NM_000169	UROD	NM_000374
GNB2L1	NM_006098	VNN3	NM_018399
GPR74	NM_004885	WBP4	NM_007187
H4F2		ZNF155	NM_003445
·	NM_003548	ZNF300	_
HAVCR-1	NM_012206	ZINF 3UU	NM_052860
HHLA2	NM_007072		
HNF4a7	AF509467		
IFNA10	NM_002171		
INSR	NM_000208	1	

(20/41)

Fig. 15A

		Direct	In vitro	Indirect	Sequence Bases	ORGANISM
Regulator	Target Gene	Reference	Reference	Reference	Reference	Organism).
HNF4a	GST-YA	and the second s	Carpenday rouse , Marketon is a . D. S. Try and the Carpenda Carpenda Carpenda Carpenda Carpenda Carpenda Carp	Paulson 1990		human
HNF4a	TTR		Sladek 1990	Sladek 1990, costa 1991		human
HNF4a	ApoC3		Sladek 1990	Sladek 1990		human
HNF4a	ApoA1		Sladek 1990	Sladek 1990		human
HNF4a	serpina		Sladek 1990	Sladek 1990		human
HNF4a HNF4a	Pktr		Sladek 1990	Sladek 1990		human
HNF4a	cyp2c13 alb		herbst 1991	herbst 1991	eguchi 1991	rat
HNF4a	tir		herbst 1991	herbst 1991		rat
HNF4a	hnf1a		HEIDST 1991	lian 1991		ral human
HNF4a	19		crossley 1991	100 1		human
HNF4a	hnfta		4,200,07	kuo 1992		human
HNF4a	apob		ladias 1992	ladias 1992		human
HNF4a	ApoC3		ladias 1992	ladias 1992		human
HNF4a	apoa2		ladias 1992	ladias 1992		human
HNF4a	pkir			puzenał 1992		human
HNF4a	(9			reljnen 1992		human
HNF4a	If			schaeffer 1993		human
HNF4a	hnf1a			zapp 1993		xenopus
HNF4a HNF4a	pck1		angrand 1994	angrand 1994		rat
HNF4a	pck2 cyp2c2		angrand 1994	angrand 1994		ral
HNF4a	cyp2c1		chen 1993 chen 1993	chen 1993		human
HNF4a	cyp2c3		chen 1993	chen 1993 chen 1993		human
HNF4a	cyp7a1		chiang 1994	chiang 1994		human
HNF4a	ApoA1		fuemkranz 1994	fuernkranz 1994		rat human
HNF4a	CEACAM1		hauck 1994	hauck 1994		human
HNF4ct	apoa4		klistaki 1994	klistaki 1994		human
HNF4a	pkir			liimatta 1994		ral
HNF4a	a2m		matthijs 1994			human
HNF4a	pkir	miquerol 1994				human
HNF4a	rbp2			nakshatri 1994		rodent
HNF4a	otc		* 400.4	nishiyori 1994		mice
HNF4a HNF4a	acox1		winraw 1994	winrow 1994		rat .
HNF4a	hsd17b4		wintow 1994	winrow 1994		rat
1011 100	17		1995	erdmann 1994, greenberg 1995		human
HNF4a	f8		figueiredo 1995	figueiredo 1995		human
HNF4a	өро		galson 1995	galson 1995		human
HNF4a	сур2с9		ibeanu 1995	ibeanu 1995		human
HNF4a	ambp		rouet 1995	rouel 1995		human
HNF4a	cyp2c23		roussel 1995			ral
HNF4a	cyp2d6		caims 1996	cairns 1996		human
HNF4cı	serpinc1		Fernandez-Rachubinski	Fernandez-Rachubinski 1996	ı	human
HNF4a	bf ·		1996	garnier 1996		human
HNF4ca	f10	·	hung 1996	hung 1996		human
HNF4cz	prir	1	moldrup 1996	moldrup 1996		rat
HNF4a	mst1		waltz 1996	wallz 1996		human
HNF4a	lipc			chang 1997		human
HNF4cc	g6pc		lin 1997	lin 1997		human
HNF4rc	SLC2A2			stoffel 1997		mouse
HNF4a	aldob			stoffel 1997		niouse
HNF4a HNF4a	gadp			stoffel 1997		mouse
HNF4a	fabpt		valeomeri 1007	stoffel 1997		mouse
HNF4a	cyp2a4 f12		yokomori 1997 farsetti 1998			mouse
HNF4a	cyp3a23		huss 1998	huss 1998		human rat
HNF4cz	shbg		janne 1998	janne 1998		rai human
HNF4a	apoc2		kardassis 1998	kardassis 1998		human
HNF4a	afp			magee 1998		human
HNF4a	HMGCS2		rodriguez 1998	rodriguez 1998		rodent
HNF4a	ALDH3A1		boesch 1999	boesch 1999		rat
HNF4cı	serplna1		hu 1 9 99	hu 1999		human
HNF4a	cyp3a1			ogino 1999		rat
HNF4a	aldh2			pinaire 1999		human
HNF4ca HNF4ca	cyp2c12 GUCY2C		sasaki 1999	sasaki 1999		ral
HNF4a	ang		swenson 1999 yanai 1999	swenson 1999 yanai 1999		human
HNF4a	ada		dusing 2000	Janua 1999		human himan
HNF4ct	hní6		lahuna 2000	lahuna 2000		human human
						191191

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Fig. 15B

TABLE S4		Direct	In with a 1.1.1 in	Indiract		
Regulator	Target Gene	Reference	In vitro Reference	Indirect Reference	Sequence Base	
HNF4a	hadhb	Training T	nicolas-frances 2000	nicolas-frances 2000	Reference	Organisi
1000 400	pax4		smith 2000	smith 2000		human
	ins		211101 2000			human
HNF4a	ogdh			wang 2000 wang 2000		mouse
HNF4ce	ucp2			_		monse
HNF4a	hni4a		haith 2004	wang 2000		mouse
HNF4ca			bailly 2001	bailly 2001		human
HNF4a	ghr		jiang 2001	jiang 2001		bovine
HNF4 α	cyp3a4			jover 2001		human
HNF4a	cyp3a5		·	Jover 2001		human
HNF4a	cyp3a6 cyp2b6			jover 2001		human
HNF4a	- ·			jover 2001		human
HNF4a	cyp2c9 fmo1	•		jover 2001 luo 2001		human
HNF4a	cyp3a16		nakarama 2001			rabbit
HNF4cz	akr1c4		nakayama2001 ozeki 2001	nakayama2001 ozeki 2001	· •	mouse
HNF4a	cyp8b1					human
HNF4α	hpd		zhang 2001	zhang 2001		human
HNF4a	cyp27		aarenstrup 2002 garuti 2002	aarenstrup 2002		rat
HNF4a	NOS2A		guo 2002	garuti 2002 guo 2002		human
HNF4α	cpt1a		900 2002	louel 2002		rat
HNF4ce	ppara		pineda-toma 2002	pineda-torra 2002	•	human
HNF4a	gk		roth 2002	pilleda-tolla 2002		human
HNF4u	Serpina1	Soutoglou 2002	1001 2002			rat
HNF1a	FGA	Outrogion 1002		baumhueter 1990	!	human
HNF1a	FGB			baumhueter 1990		
HNF1a	FGG			baumhueter 1990		
HNF1a	afp			baumhueter 1990		
HNF1a	serpina1			baumhueler 1990		•
HNF1a	afm			herbst 1991	carachini 1000	ant thanh
HNF1 α	alm			tronche 1991	cereghini 1990	rat (herb
HNF1a			gonzalez 1990, hayashi	HOURIS 1331	î	rat
	cyp2e1		1991			enimal
HNF1a	aldob		raymondjean 1991			rat
HNF1a	aldob		ito 1990		_	rat
HNF1a	inflored			suwanichkul 1990, babajko	•	
	iglbp1			1993		human
HNF1a	igfbp1			powell 1993		human
HNF1a	igfbp1			suh 1995, suh 1997		rat
	сгр			Ioniatti 1990		
$HNF1\alpha$	apoa2			chambaz 1991		human
HNF1a	ltr	,		costa 1991		mouse
HNF1cx	lt			herbst 1991		rat
HNF1a	hdlbp				drewes 1991	xenopus
HNF1a	rbp5			tripodi 1991		human
HNF1a	f2		bancroft 1992	bancroft 1992		human
HNF1a	apob		brooks 1992			human
HNF1a	insr		cameron 1992			human
HNF1a	insr		cameron 1992			human
HNF1a	agt			congiu 1992		mouse
HNF1a	ins			emens 1992		rat
HNF1a	pklr		puzenal 1992			
HNF1a HNF1a	tat		schweizer-groyer 1992	#URAGOS 4000 F.1.1		rat
BIAL RX	siat1		svensson 1992	svensson 1992, bois-joyeux	•	human
HNF1a	adh1			1995 van ooii 1992		
HNF1a	cthbp			van ooji 1992	hohan 1002	human
HNF1ct	afp			bernier 1993	behan 1993	human
HNF1a	fgb		dalmon 1993	dalmon 1993		human
HNF1a	lyz		COUNTY 1999	Agillion 1999	araine 1002	human
HNF1a	aldob			araani 1002	grajer 1993	chicken
HNF1a	lbg			gregori 1993 hayashi 1993		human
HNF1cc	apoa1			krilis 1993		human
HNF1a	apoc3			kritis 1993		
HNF1a	ctb		li 1996	ku 1993, li 1996		mouse
HNF1a	fgb		1) TOUT	roberts 1993		mouse
HNF1a	btoc			berg 1994		xenopus
HNF1a	serpina1			bulla 1994		human
HNF1a	gsla2		clairmont 1994	-unu IVVT		human
	cyp2c13		TOTAL PARTY OF THE	legraverend 1994		human
HNF1α	pkir	migueral 1994		iograficity 1007		human
HNF1a	aubeb	,	olsen 1994	olsen 1994		human
HNF1a	si		vw 1994	พบ 1994		human
			1 # # T	THE THET		HOITIGHT

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Fig. 15C

		Di				
TABLE S4	/ Tamel Good Till	Direct	In vitro	Indirect Reference	Sequence Based	
HIVETU	Target Gene CABPA	- Maisialiney 1 1 1	-: Reference - No. 1995		Keleleuce	
HNF1a	FGA			arenzana 1995 hu 1995		human human
HNF1a	igf1			kutik 1995, nolten 1995		salmon, human
HNFta	cyp2e1		līu 1995	ilu 1995, lerche 1996		taj
HNF1a	ambp		rouet 1995	rouet 1995		human
HNF1a	ddc		aguanno 1996	aguanno 1996		human
HNF1a	18		mcglynn 1996	mcglynn 1996		human
HNF1a	plg		meroni 1996	meroni 1996		human
HNF1a	pah			pontoglio 1996		mouse
HNF1a	hmgcs2				boukaflane 1997	human
HNF1a	lipc			chang 1997		ral
HNF1a HNF1a	cyp2h1		dogra 1997	dogra 1997		chicken
HNF1a	ugt2b1 guanylin		hansen 1997	hansen 1997		human, rat
HNF1a	36b		hochman 1997 lin 1997	hochman 1997 Im 1997		mouse
HNF1a	cyp2e1		McGehee 1997	McGehee 1997		human rodent
HNF1cc	pah		Pontoglio 1997	moderice 1991		Monze
HNF.1a	ipal		Taylor 1997			mouse
HNF1a	hnf4a		•	bailly 1998		ral
HNF1a	hnf3a			bailly 1998		rai
HNF1a	cebpa			bailly 1998		raì
HNF1a	g6pc		lin 1999	lin 1998		human
HNF1a	alp		magee 1998	magee 1998		human
HNF1a	SLC5A1		rhoads 1998			rat
HNF1a	si		I maa	rodolosse 1998		human
HNF1ce HNF1ce	gc CIII TOA4		song y 1998	song y 1998		human
HNF1a	SULT2A1		song c 1998	song c 1998		ral
HNF1a	proc g6pc		streeper 1998	spek 1998		human
HNF1a	SLC10A1		trauner 1998	streeper 1998		human human
HNF1ct	ins		1000	wang 1998		human human
HNF1a	ugtiai		bemard 1999	Tang 1000		human, mouse
HNF1a	cyp7a1		chen 1999			human
HNF1a	dpp6			erickson 1999		human
HNF1a	serpina6		hu 1999	hu 1999		human
HNF1a	igf1			meton 1999		salmon
HNF1a	ins		okila 1999	okita 1999		human
HNF1cz	CYP27A1		rao 1999	rao 1999		ral
HNF1a HNF1a	lct SI CEA1		spodsberg 1999	spodsberg 1999		mice
HNF1a	SLC5A1 fabp1			wood 1999		human
HNF1a	сур7а1		antes 2000	akiyama 2000 antes 2000		mouse
HNF1a	slc2a2		cha 2000	cha 2000		mice human
HNF1a	dpp6		erickson 2000	erickson 2000		human human
HNF1a	UGT2B17		gregory 2000	gregory 2000		human
HNF1a	UGT2B7		ishii 2000	ishii 2000		numan
HNF1a	ugt1a7		metz 2000	metz 2000		rat
HNF1a	fech			muppala 2000	1	nouse
HNF1a	gjb1		piechocki 2000	piechocki 2000	1	numan
HNF1a	SLC5A2		Pontoglio 2000	pontoglio 2000		numan
HNF1ce HNF1ce	pax4		smith 2000	smith 2000		numan
HNF1a	ogdh aldob			wang 2000 wang 2000		rat
HNF1a	ins			wang 2000		rat 1
HNF1a	SLC5A2			wang 2000		al al
HNF1a	pklr			wang 2000		at
HNF1a	hmgar			wang 2000		al al
HNF1a	hnf4a		bailly 2001	bailly 2001		uman
HNF1a	pdx1		ben-shushan 2001	ben-shushan 2001		ruman
HNF1a	hnf4a7	Boj 2001				nouse
—	hnf3g	Boj 2001			ſ	nouse
	hn[4g	Boj 2001		•	1	nouse
_	gck		cha 2001	cha 2001		iuman
	hnf4a	Hatzis 2001	hatzis 2001	hatzis 2001		luman
	g6pc c6pti			hiraiwa 2001		nouse
	g6pt1 slc21a6		iuno 2001	hiraiwa 2001		nouse
	slc21a8		jung 2001	jung 2001 jung 2001		iuman
	ngn3			lee 2001	_	ເບກາລາ ເບກາລາ
	igfbp1			leu 2001		odent
	g6p	•		leu 2001		odent
HNF1a	alp			leu 2001		odent
	fmo1			luo 2001		abbil, human

Fig. 15D

TABLE S		Direct	In vitro	Indirect	Continue Paradia Paradia Continue
Regulator	Target Gene		Reference	THE PROPERTY OF THE PROPERTY O	Sequence Based Organism
HNF1a	CYP27A1	A TELEVISION & TELEVISION AND ADDRESS OF THE PERSON ADDRESS OF THE PERSON AND ADDRESS OF THE PERSON AND ADDRESS OF THE PERSON	memom 2001	ma management and and the state of the state	hamster
HNF1a	AKR1C4		oxeki 2001	ozeki 2001	human
HNF1a	NR5A2		pare 2001	pare 2001	mouse
HNF1a	cyp2c11		park 2001	park 2001	rodent
HNF1a	cyp2a2		park 2001	park 2001	rodent
$HNF1\alpha$	cyp4a2		park 2001	park 2001	rodent
HNF1ca	pklr		parrizas 2001	•	human
HNF1a	slc2a2		parrizas 2001		human
HNF1a	pah		parrizas 2001		human
$HNF1\alpha$	c8a		•	pontoglio 2001	mouse
HNF1a	c 5			pontoglio 2001	mouse
HNF1a	cyp2e1		roe 2001	,	rat
HNF1a	nr1h4		shih 2001	shìh 2001	mouse
HNF1a	SLC10A2		shih 2001	shih 2001	mouse
HNF1a	SLC17A1			soumounou 2001	human, mouse
HNF1a	hnf4a7			thomas 2001	human
HNF1a	ins			yamakawa 2001	human
HNF1a	Nr5a2			zhang 2001	naman
HNF1a	SLC5A1			vayro 2001	sheep
HNF1a	slc2a2		ban 2002	ban 2002	human
HNF1a	si			boudreau 2002	mouse
HNF1a	SLC17A1			cheret 2002	mouse
$HNF1\alpha$	SLC10A1		geier 2002		rat
HNF1cc	UGT2B17		gregory 2002	gregory 2002	human
HNF1α	hnf4a7			hansen 2002	mouse
HNF1a	gjb1			koffler 2002	rat
HNF1a	AKR1C4		ozeki 2002	ozeki 2002	human
HNF1a	cldn2			sakaguchi 2002	human, mouse
HNF1a	fgfr4		shah 2002	shah 2002	human
HNF1a	igf1		•	yang 2002	human/rat
HNF1a	mif			yang 2002	human/rat
HNF1α	Serpina1	Soutoglou 2002			human
HNF1α	c1		zahedi 2002		human

Fig. 16

A1BG	RefSeq 17.			SENDING SERVICE			ReiSeg		'ReiSeq 🤼
	NM_130786	DHFR	NM_000791	GSS	NM_000178	ORC1L	NM_004153	UGT1A1	NM_000463
AASS	NM_005763	DKFZP434J037	NM_030952	H3FF	NM_003533	PABPC1	NM_002568		NM_001073
ABCA8	NM_007168	DKFZP564O0523		H4FK	NM_003546	PCDHA12	NM_018903		NM_001076
ABCB11	NM_003742	DKFZP586A0522		HABP2	NM_004132	PCK1	NM_002591	URKL1	NM_017859
ABCC2	NM_000392	DXF68S1E	NM_012080	HBP1	NM_012257	PHTF1	NM_006608	VCP	NM_007126
ABL2	NM_007314	E2F1	NM_005225	HCAP-G	NM_022346	PIK4CB	NM_002651	VTN	NM_000638
ACVR1	NM_001105	E2F1	NM_005225	HESX1	NM_003865	PLGL	NM_002665	WDR12	NM_018256
ADH1A	NM_000667	ElF4A1	NM_001416	HIVEP3	NM_024503	POLR2D	NM_004805	WDR5B	NM_019069
ADH1B	NM_000668	EIF4E	NM_001968	HMGCR	NM <u>·</u> 000859	POLS	NM_006999		_
AF038169	NM_013310	ELOVL1	NM_016031	HNF4a7	AF509467	PON1	NM_000446		
AGTR1	NM_000685	EPHA1	NM_005232	HNMT	NM_006895	PPFIA1	NM_003626		
AKR1C4	NM_001818	F11	NM_019559	HNRPR	NM_005826	PPP2R5A	NM_006243	Î	
ALDH3A1	NM_000691	F9	NM_000133	HSD17B4	NM_000414	PRO1855	NM_018509		
ALDH5A1	NM_001080	FABP5	NM_001444	HSP105B	NM_006644	PSMA1	NM_002786		
AMBP	NM_001633	FACTP140	NM_007192	HSPA1B	NM_005346	PSMB1	NM_002793		
AMT	NM_000481	FADS3	NM_021727	HTR2B	NM_000867	PTPRR	NM_002849		
APCS	NM_001639	FLJ10209	NM_018026	IF.	NM_000204	REA	NM_007273	Ι,	
APOH	NM_000042	FLJ10407	NM_018087	INSM2	NM_032594	RING1	NM_002931		
ASPA	NM_000049	FLJ10415	NM_018089	IRF3	NM_001571	RNF20	NM_019592		
BCAR1	NM_014567	FLJ10578	NM_018144	IRF6	NM_006147	RPL35	NM_007209		
BCKDHA	NM_000709	FLJ10650	NM_018168	ITGAV	NM_002210	RPL37AP1			
BF	NM_001710	FLJ11029	NM_018304	ITIH1	NM_002210	•	NG_000988		•
BM039	NM_018455	FLJ11105	NM_018323	JIK	. -	RPLP1	NM_001003		
BNIP3L	NM_004331	FLJ11301	NM_018385		NM_016281	RPS6KA5	NM_004755		
BTN3A2	NM_007047	FLJ11726	NM_024971	KIAA0806	NM_014813	RRP46	NM_020158		
C1S	NM_001734	FLJ11773	-	KIAA10EC	NM_014940	SART3	NM_014706		
C2	NM_000063	FLJ12552	NM_021934	KIAA1056	NM_014894	SAS10	NM_020368		
	-	1	NM_022832	KLF3	NM_016531	SCYB13	NM_006419		
	NM_015638	L	NM_032174	LIMK1	NM_016735	SEC10L1	NM_006544		
C8B	•		NM_024573	LOC51060	NM_015913	SERPING1	_		
C8G	NM_000606		NM_024773	LOC51074	NM_015957	SERPINI1	NM_005025		
	NM_000720	1	NM_022736	LOC51287	NM_016565	SILV	NM_006928		
CASP2	NM_032982	F .	NM_017659	LOC51633	NM_016023	SLC1A3	NM_004172		
	. -	I	NM_017691	LOC51646	NM_016061	SLC25A13	NM_014251	:	
			NM_017814	LOC56906	NM_020147	SLC7A9	NM_014270		
CDC2L5		l	NM_017909	LOC81558	NM_030802	SMARCC1	NM_003074		
			NM_017924	LOH11CR2A	NM_014622	SMCY	NM_004653		•
	-		NM_017956	M17S2	NM_031858	SNRPD2	NM_004597		
	NM_005507	FLJ21934	NM_024743	MAP2K5	NM_002757	SNW1	NM_012245		
CH25H	NM_003956	FLJ21963	NM_024560	MGC10500	NM_031477	SNX3	NM_003795		
CLCN3	NM_001829	1	NM_024085	MGC13053	NM_032710	SPG4	NM_014946		
			NM_024713	MGC16169	NM_033115	SPINK1	NM_003122		
	-		NM_025192	MGC16386	NM_080668	SPP2	NM_006944		
	_		NM_025115	MGC4189	NM_032308	SRF	NM_003131		
	-		NM_024956	MGST3	NM_004528	STMN2	NM_007029		
	_		NM_022761	MN1	NM_002430	TAF2GL	NG_001012		
_			NM_024783	NEK6	NM_014397	TAT	NM_000353		
			NM_022006	NFKBIA	NM_020529	TBX2	_		
			NM_000151	NFKBIA	NM_020529	TCEB3	NM_005994		
			NM_002040	NFKBIA			NM_003198		
					NM_020529	TM4SF4	NM_004617		
	NM_001891		NM_033036	NOLC1	NM_004741	TMF1	NM_007114		
	NM_134268		NM_004193	NR112		TMOD2	NM_014548		
				NTF2			NM_000043		
	NM_000786		NM_002086	OAT	NM_000274		NM_003810		
	NM_005800		NM_001511	OAZ2			NM_014820		
DB2	NM_000107	GRO3	NM_002090	JOGFR	NM_007346	TSG101	NM_006292		

Fig. 17

Name -	RefSeq	Name	RefSeq 18	Name 1	RefSeq. (4. 50	Name .	RefSeq
AASS	NM_005763	FLJ11271	NM_018373	JIK	NM_016281	SEMA6A	NM_020796
ABCB8	NM_007188	FLJ11301	NM_018385	KIAA0660	NM_012297	SERPINB8	-
ACPP	NM_001099	FLJ11773	NM_021934	KIAA0712	NM_014715	SERPING1	NM_000062
ACVR1	NM_001105	FLJ12770	NM_032174	KIAA0806	NM_014813	SERPINI1	NM_005025
ADH1A	NM_000667	FLJ12910	NM_024573	KIAA0872	NM_014940	SH3BGRL	NM_003022
AF038169	NM_013310	FLJ13220	NM_021927	KIAA1056	NM_014894	SLC1A3	NM_004172
AF15Q14	NM_020380	FLJ13798	NM_024773	KRTAP1.1	NM_030967	SNRPD2	NM_004597
AGT	NM_000029	FLJ13955	NM_024759	LAMC2	NM_018891	SNW1	NM_012245
AMBP	NM_001633	FLJ14153	NM_022736	LBC	NM_006738	SPG4	NM_014946
AMT	NM_000481	FLJ14486	NM_032792	LOC51060	NM_015913	SPINK1	NM_003122
APCS	NM_001639	FLJ20084	NM_017659	LOC51287	NM_016565	TEGT	NM_003217
APOH	NM_000042	FLJ20156	NM_017691	LOC51633	NM_016023	TMF1	NM_007114
ARL1	NM_001177	FLJ20422	NM_017814 ·	LOC56906	NM_020147	TNFRSF6	NM_000043
BBP	NM_032027	FLJ20627	NM_017909	LOC81558	NM_030802	TNFRSF6	NM_000043
BCKDHA	NM_000709	FLJ20643	NM_017916	LOH11CR2A		TNFRSF6	NM_000043
BF	NM_001710	FLJ20671	NM_017924	LUC7A	NM_016424	TNFRSF6	NM_000043
BTN3A2	NM_007047	FLJ20772	NM_017956	MDH1	NM_005917	TNFSF10	NM_003810
C1\$	NM_001734	FLJ21272	NM_025032	MDS029	NM_018464	TOMM70A	NM_014820
C20orf188	NM_015638	FLJ21934	NM_024743	MEIS1	NM_002398	UGT2B15	NM_001076
C2F	NM_006331	FLJ21963	NM_024560	MGC13040	NM_032930	UGT2B17	NM_001077
C8orf4	NM_020130	FLJ22169	NM_024085	MGC13053	NM_032710	VCP	NM_007126
CCT8	NM_006585	FLJ23263	NM_025115	MGC19595	NM_033415	VTN	NM_000638
CDC2L5	NM_003718	FLJ23375	NM_024956	MGC3020	NM_024048	WDR12	NM_018256
CH25H	NM_003956	GABARAPL1	NM_031412	MGC3413	NM_032678		NM_020933
CIR	NM_004882	GABPA	NM_002040	MGC4189	NM_032308		
CLCN4	NM_001830	GCP3	NM_006322	MGST3	NM_004528		
CLDN2	NM_020384	GJB1	NM_000166	MTERF	NM_006980		
CLLD8	NM_031915	GLA	NM_000169	NET-6	NM_014399		
CLNS1A	NM_001293	GRB2	NM_002086	NOLC1	NM_004741		
CLONE24922	NM_015679	GRO1	NM_001511	NOVA1	NM_006489		
CMG1	NM_025103	GRO3	NM_002090	NR0B2	NM_021969		
COPB2	NM_004766	GSS	NM_000178	NUDT2	NM_001161		
COPS7A	NM_016319	GSTA4	NM_001512	OGFR	NM_007346		
COX4I1	NM_001861	GTF2E1	NM_005513	ORC1L	NM_004153		
COX7A2L	NM_004718	H4FA	NM_003538	PAPA-1	NM_031288		
CRI1	NM_014335	H4FH	NM_003543	PEX6	NM_000287		
CSN2	NM_001891	HABP2	NM_004132	PMAIP1	NM_021127		
CYP3A43	NM_022820	HASJ4442	NM_017528	PPFIA1	NM_003626		
DKFZp761D221	_	HBOA	NM_007067	PPFIBP1	NM_003622		
DKFZp761J139	NM_032280	HBP1	NM_012257	PPP1R3D	NM_006242		
EED	NM_003797	HLA-G	NM_002127	PSMA1	NM_002786		
EGR2	NM_000399	HMG2	NM_002129	PSMB1	NM_002793		
EHD4	NM_014599	HNF4a7	AF509467	PTPRN2	NM_002847		
EHF	NM_012153	HNRPA2B1	NM_031243	REA	NM_007273	•	
EIF4E	NM_001968	HNRPR	NM_005826	RECK	NM_021111		
F11	NM_019559	HSD17B4	NM_000414	RIG-I	NM_014314		
F2RL2	NM_004101	HSN44A4A	NM_015372	RPC32	NM_006467		
FABP5	NM_001444	HSP105B	NM_006644	RPL36P1	NG_000983		
FER1L3	NM_133337	HSPA1B	NM_005346 '	RPS6KA5	NM_004755		
FLJ10342	NM_018064	HSPC125	NM_014165	RRP46	NM_020158		
FLJ10407	NM_018087	HT007	NM_018480	SAMHD1	NM_015474		
FLJ10415		HTR2B	NM_000867	SART3	NM_014706		
FLJ10482	NM_018107	humNRDR	NM_021004	SAS10	NM_020368		
FLJ10650	NM_018168	IGSF3	NM_001542		NM_019846		
FLJ11029	NM_018304	IRF3	NM_001571	SEC10L1	NM_006544		

(26/41)

Fig. 18A

24172			Name & RelSequit		ReiSeq 3	Gene Name	ReiSeq 5	Gena)lam	e Reiseq7.	Gana Nam ReiSeq	S-Gene Nam	Refser :::
Section Sect			A5 NM 052968 B NM 000384		NM_005768 NM_017546	ICPT1B	NM 004377	DNAJA3	NM 005147	FLJ11184 NM 018352	FLJ22028	NIA 024854
ABSC M. 0.0376	4TM NM_02047	020470 APO	C2 NM_000483	C4A	NM_007293	CRADD	NM_003805	DNAJB4	NIA 007034	IFLJ11198 NM 018358	FLJ22191	NM 024085 NM 025231
ABOCS MR 014458 MR 01455 Ceris MR 04455 Ceris MR	ASS NH 00576					ICREBL2 ICRFG	NM_001310 NM_012341		NM_012328 NM_005851	IFUJ11274 NK 018375	FLJ22353	NM_024587
ABCRIT M 000392	B026190 NW 01445	014458 AQP	3 NM 004925	C6cri11	NM 005452	CRI1	NM 014335	DPAGT1	NM 001382	IFU11342 NM 018394	FLJ22501	NM 024747
ABCC2 NN 000392	BCB10 NM 01208	012089 AQP	9 NM_020980	C7crf10		CRKL				IFLJ11526 NM 024632 IFLJ11728 NM 024971	FLJ22551	NM_024708 NM_024520
ABCC MM, 001716 ABCC MM, 001717 C66rd MM, 202130 CREP3 MM, 001620 DUSPE MM, 022522 FL11184B MM, 202135 FL122849 MM, 202134 ABCC MM, 002491 ABC MM, 0				C8B C8G	830000, MM	CROT	NM_021151	DUSP11	NM 003584	IFW11767 NM 024593	FLJ22557	NM 024713
ABCE MI, 002945 ART MI, 002945 ART MI, 002947 CACAR2D MI, 002950 CRY1 MI, 002950 CRY2 MI, 0029	BCC3 NM_003786	.003788 ARG	2 NM 001172	C8orf4	NM_020130	ICRSP3		DUSP6		IFLJ11838 NM 024664 IFLJ11848 NM 025155	FLJ2257B FLJ22637	NM 024864 NM_025165
ABCGS NM 002437 ARL5 NM 001177 CACNA202 NM 006500 CPCY NM 006889 EFG1 NM 024696 FL12233 NM 023027 FL122272 NM 012462 CS NM 004077 CARD15 NM 007270 CARD15	3CC6 NM_00117 BCE1 NM_00294					CRSP9	NM 004270		NM 004714	[FLJ12171 NM_024619	FLJ22649	NM 021928
ABLIA M 060727 ARLS MW 02239 CARACU MN 09167 CARSP M 05177 CARSP M 02392 CARACU MN 09168 CARS MN 09167 CARSP M 02392 CARACU MN 09168 CARS MN 09167 CARSP M 02392 CARACU MN 09168 CARS MN 09167 CARSP M 02392 CARACU MN 09168 CARS MN 09167 CARSP M 02392 CARACU MN 09168 CARS MN 09167 CARSP M 02392 CAR	BCG1 NM 004919	004915 ARL1	NM 001177	CACNA2D2	NM 006030	CRYZ	NM 001889	EFG1	NM 024996	FLJ12439 NM 023077	FLJ22729	NM 024683
ASS NM, 016222 ARPCS NM 005717 CASP2 NM 023982 (CSNR2A) NM 001699 (CSNR2A) NM 001699 (CSNR2A) NM 001691 (ASS NM 001690) (CSNR2A) NM 001691 (ASS NM 001690) (CSNR2A) NM 001691 (ASS NM 001690) (CSNR2A) NM 001691 (ASS NM 001691) (ASS NM 00169	BLIM NM 006720		NM 005737	CAMK2D CARD15		CS CSDUFD1				FLJ12552 NM_022832	FLJ22865	NM, 025109
ACADS M. 00811 ASS3 NN. 01611 CATSER NN. 02583 CSTF1 NN. 001226 EFESS NN. 00168 FLIZZES NN. 022492 FLIZZES NN. 02684 ACADV NN. 001691 ASGRI NN. 001671 CATSPER NN. 0053054 CSTF3 NN. 001326 EFESS NN. 001686 FLIZZES NN. 02684 FLIZZES NN. 02684 ACADV NN. 001696 AFF2 NN. 001686 CSS NN. 000677 CTMP NN. 001695 EFF4SEP2 NN. 001686 FLIZZES NN. 02687 ACLY NN. 001696 AFF2 NN. 001696 GSS NN. 000777 CTMP NN. 001695 EFF4SEP2 NN. 001686 FLIZZES NN. 02687 NN	BS NM_016221	_016222 ARP(C5 NM 005717	CASP2	NM 032982	CSNK2A1	NM 001895	EHM2	NM 019114	FLJ12707 NM, 022067	FLJ23071	NM 025192
ACADAY M 001609 ASGR! NN 001601 CATSPER NM 05055 CSTF3 NM 001355 EFFEE NM 001668 FL/32355 NM 02167 CTSP NM 001355 EFFE NM 001668 FL/32355 NM 022461 ACCY NM 001675 ACF NM 001675 ACF NM 001675 ACF NM 001675 ACF NM 001675 ACCY NM 001686 CBS/3 NM 000277 CTSZ MM 001355 EFF NM 001688 FL/32353 NM 022461 ACCY NM 001696 ACF NM 001696 CBS/3 NM 00277 CTSZ MM 001359 EFF NM 001688 FL/32353 NM 022461 ACCY NM 001698 ATF NM 006665 CBS/3 NM 001211 CYB5 NM 001814 EFF NM 001643 ACCY NM 001674 ACCY ACCY NM 001674 ACCY ACCY NM 001674 ACCY AC	CAA2 NM 008111	008111 ASB	NM 016115		NM_025263	CSTF1		EIF2S1 EIF2S3	NM_004094 NM_001415			NM_024643
ACLY NII. 001098 ATF4 NO 01676 EB33 MM 002707 CTSZ MT 001333 EF65 NIII. 001676 EB33 MM 002707 ACO24 NII. 001098 ATF4 NII. 006856 CB35 NIM. 002707 CTSZ NII. 001391 EL73 NII. 002403 FL/3108 NII. 002403 FL/3108 NII. 002403 ACO24 NII. 001098 ATF7 NII. 006856 CB35 NIM. 002102 CCB3 NII. 001020 NII. 001020 ATM NII. 001051 CCMG NII. 004035 CCB3 NII. 001020 CCB3 NII. 001030 CCB3 NII. 001030 ACO24 NII. 004035 ATM NII. 002501 CCMG NII. 004035 ACO24 NII. 004035 ATM NII. 002501 CCMG NII. 004035 ACO24 NII. 004035 ATM NII. 002501 CCMG NII. 004035 ACO24 NII. 004035 ATM NII. 002501 CCMG NII. 004035 ACO24 NII. 004035 ATM NII. 002501 CCMG NII. 004035 ACO24 NII. 004035 AC					NM_053054	CSTF3	NM_001326	EJF4E	NM, 001968	[FLJ12886 NM 019108	FLJ23251	NM 024818
ACOZ MM. 001998 ATFF MM. 001996 ATFF MM. 0019675 CBXS MM. 0027276 CU12 MM. 003991 ELP3 NM. 004935 ATFF MM. 009605 CBXS MM. 001707 CV95 MM. 001904 ELP2 MM. 018255 LP313168 NM. 025002 FL12343168 MM. 02604 ATFF MM. 001905 CCNG2 MM. 004060 CVP1A2 MM. 0035079 ENC1 MM. 004035 ATFM MM. 005010 ATFF MM. 005010	CF NM_014576	014576 ATF2	NM 001880	CBS	NM_000071	CTSZ						NM 025115 NM 025059
ACOXI			NM 001675 NM 006856			CUL2	NM 003591	ELF3	NIM_004433	FLJ13102 NM_024867	FLJ23441	NM_024678
ACP2 NM 001613 AFP5F1 NM 001659 CCNH NM 001659 ACTGA NM 001659 CCNH NM 001659 ACTGA NM 001659	COX1 NM_004035	.004035 ATM	NM 000051	CCNG1	NM_004060	CY85-M	NPA_030579	ENC1	NM 003633	FLJ13162 NM_025002	FLJ23408	NM 024029 NM 022761
ACTR1	CP2 NM 001610					CYP1A2	NM 000761 NM 000104	EPB72	NM 004099 NM 004431	FLJ13181 NM_025188		NIA 024725
ACTR3	CTA2 NM, 001613	001613 ATP5	G3 NM 001689	CCT6A	NM_001762	CYP21A2	NM_000500	EPI64	NM 031937	FLJ13195 NW 022906	FMR1	NIA_002024
ACVR1	CTR3 NM 005721	.005721 ATP6	G1 NL4 004888	CD68		CYP255 CYP2C8		JERBB2IP JERBB3				NM_002027 NM_002028
AD022	VR1 NM 001105	001105 ATP6	L NM 001694	CDC14A	NM 001785	CYP2D6	NM 000106	ERCC5	NM 000123	IFLJ13291 NM 032178	FOSL2	NM 005253
ADD34 NM 031480 NM 032270 ATPW NM 015684 CDC51 NM 001253 CYP3A5 NM 022820 EVC NM 014556 FLJ13611 NM 024941 FTHFD NM 012 NM 022820 EVC NM 014556 FLJ13611 NM 024941 FTHFD NM 012 NM 022820 EVC NM 014556 FLJ13611 NM 024941 FTHFD NM 012 NM 024941 FTHFD NM 024941 FTHFD NM 012 NM 024941 FTHFD NM 024944	0022 NM 016814	016814 ATP6	S14 NM 004231	CDC25A	NM_001789	CYP2E	NM_000773	ERO1L	NM_014584	FLJ13448 NM_025147	FRK	NM_004477 NM_002031
AD24		.031480 ATP7 .032270 ATPV				CYP2J2			NM 005797	[FLJ13491 NM_024623	FSTL3	NM_005660
ADHE NM 000672 B29 NM 031939 CDK13 NM 016508 CYP4F2 NM 001082 F10 NM 000504 F113769 NM 025012 FXP07 NM 025012	024 NM_022451	022451 AUP1	NM 012103	CDCA1	NM 031423	CYP3A5	NM,000777	EVG1	NM 032561	FLJ13615 NM_025114	FTSJ1	NM 012190 NM 012280
ADPRTL										FLJ13660 NNL 025197		NM_003902
ADPRTL3 NM 005485	DPRH NM 001125	001125 B3GA	T1 NM 018644	CDKN1B	NM 004064	CYP4F3	NM 000896	F12	NM 000505	FLJ13708 NM 024773	FZD1	NM 003505
AF093680 NM 013242 BAL NM 031458 CDSN NM 001624 D123 NM 006023 FAP48 NM 053274 FLJ13964 NM 032186 G3A NM 018464 CDSN NM 006023 FAP48 NM 053274 FLJ13964 NM 032186 G3A NM 018464 CDSN NM 006023 FAP48 NM 053274 FLJ13964 NM 032186 G3A NM 018464 CDSN NM 006023 FAP48 NM 053274 FLJ13964 NM 032186 G3A NM 018464 CDSN NM 006023 FAP48 NM 032639 FLJ14153 NM 022736 G6PC NM 006024 NM 006023 FAP2 NM 032639 FLJ14153 NM 022736 G6PC NM 006024 NM 006023 FAP2 NM 012304 GAPD NM 012304 FAP2 NM 012304 FAP2 NM 012304 GAPD NM 012304 FAP2 NM 012304 FAP	PRTL3 NM 005485	005485 BACE	NM 012104	CDKN1B	NM 004064	CYP8B1				FLJ13949 NM, 025077 FLJ13952 NM 024798		NM, 017412 NM 015714
AF140225 NM 030799 BAT1 NM 004640 CDW92 NM 080546 D13S106E NM 005800 FAPP2 NM 032639 FLJ14153 NM 022736 G6PC NM 0006411 NM 000027 BAT4 NM 01348 CERD4 NM 012074 DAG1 NM 004599 BAZ1A NM 01348 CERD4 NM 012074 DAG1 NM 004393 FBX04 NM 012172 FLJ14431 NM 032783 GAB1 NM 002783 GAB1 NM 005800 FAPP2 NM 012172 FLJ14431 NM 032783 GAB1 NM 002783 GAB1 NM 005801 NM 00574 FBX024 NM 012172 FLJ14431 NM 032783 GAB1 NM 002783 GAB1 NM 002777 FLJ1462 NM 032818 GAB1 NM 002777 FLJ1462 NM 032848 GCP NM 003842 GAB1 NM 002777 FLJ1482 NM 032848 GCP NM 003841 NM 002785 GCP NM 00						Cyl19	NM, 020682	FACTP140	NM_007192	FLJ13962 NM 024862	G10	NM_003910
AGA NM 000027 BAT4 NM 03177 CEP3 NM 006449 DAF NM 00574 FBX024 NM 012172 FLJ14393 NM 032778 GAB1 NM 002783 GAB2 NM 012176 FLJ14511 NM 033087 GAB2 NM 002783 GAB2 NM 002785	140225 NM 030799	030799 BAT1	NM 004640	CDW92	NM 080546	D13S106E	NM_005800	FAPP2	NM_032639	FLJ14153 NM_022736		NM_019101 NM_000151
AGM1 NM 015599 BAZ1A NM 013448 CERD4 NM 012074 DAG1 NM 004393 FBXO4 NM 012176 FLJ14511 NM 033087 GABPA NM 002411 BAZ1B NM 032408 CETN2 NM 004344 DBI NM 020548 FBXO8 NM 012180 FLJ14621 NM 032811 GABPB2 NM 002 AGXT2 NM 001190 CEZANNE NM 020205 DBP NM 001352 FBXW2 NM 012164 FLJ14624 NM 032813 GADD45G NM 006 AGXT2 NM 031930 CC005 NM 014887 DC11 NM 031279 BCD02 NM 031930 CG005 NM 014887 DC11 NM 020186 FDXR NM 024417 FLJ14681 NM 032824 GBE1 NM 006 AK2 NM 001625 BCS1L NM 004328 CGF01 NM 015935 DC8 NM 015471 FEM1A NM 020177 FLJ14697 NM 032848 GCFR NM 005 AK2 NM 001625 BCS1L NM 004328 CGF01 NM 015941 DCK NM 000788 FEM1B NM 015322 FLJ14840 NM 032850 GCFR NM 005 AKAP13 NM 007200 BET1 NM 005868 CGF11 NM 015941 DCK NM 000788 FEM1B NM 015322 FLJ14840 NM 032850 GCFR NM 005												NM_001467
AGT NM 000029 BCAT2 NM 00190 CEZANNE NM 020205 DBP NM 001352 FBXW2 NM 012164 FLJ14624 NM 032813 GADD45G NM 006 AGXT2 NM 031900 BCCIP NM 016567 CFL2 NM 021914 DBT NM 001918 FDX1 NM 004109 FLJ14642 NM 032818 GAPD NM 006 AGXT2L1 NM 031279 BCD02 NM 031938 CG005 NM 014887 DC11 NM 020186 FDXR NM 024417 FLJ14681 NM 032824 GBE1 NM 006 AK2 NM 001622 BCL6 NM 001706 CGBP NM 014593 DC13 NM 020188 FE65L2 NM 006051 FLJ14697 NM 032826 GC20 NM 005 AK2 NM 001625 BCS1L NM 004328 CGI-01 NM 015935 DC8 NM 015471 FEM1A NM 020177 FLJ14827 NM 032848 GCHFR NM 005 AKAP13 NM 007200 BET1 NM 005868 CGI-11 NM 015941 DCK NM 000788 FEM1B NM 015322 FLJ14840 NM 032850 GCKR NM 0015	SM1 NM 015599	015599 BAZ1	A NM 013448	CERD4	NM_012074	DAG1	NM_004393	FBXO4	NM 012176	FLJ14511 NM_033087	GABPA	NM_002040
AGXT2 NM_031900 BCCIP NM 016567 CFL2 NM 021914 DBT NM_001918 FDX1 NM_004109 FLJ14642 NM_032818 GAPD NM_002 AGXT2L1 NM 031279 BCD02 NM 031938 CG005 NM 014887 DC11 NM_020186 FDXR NM_024417 FLJ14681 NM_032824 GBE1 NM_000 AHSG NM_001622 BCL6 NM_001706 CGBP NM_014593 DC13 NM_020188 FE65L2 NM_006051 FLJ14697 NM_032826 GC20 NM_005 AK2 NM_001625 BCS1L NM_004328 CGI-01 NM_015935 DC8 NM_015471 FEM1A NM_020177 FLJ14827 NM_032848 GCHFR NM_005 AKAP13 NM_007200 BET1 NM_005868 CGI-11 NM_015941 DCK NM_000788 FEM1B NM_015322 FLJ14840 NM_032850 GCKR NM_001	ST NM 000029	000029 BCAT	2 NM 001190	CEZANNE		DBP				JFLJ14621 NM 032811 JFLJ14624 NM 032813		NM 002041 NM 006705
AHSG NM 001622 BCL6 NM 001706 CGBP NM 014593 DC13 NM 020188 FE65L2 NM 006051 FLJ14697 NM 032826 GC20 NM 005 NM 01625 BCS1L NM 004328 CGF01 NM 015935 DC8 NM 015471 FEM1A NM 020177 FLJ14827 NM 032848 GCHFR NM 005 NM 007200 BET1 NM 005868 CGF11 NM 015941 DCK NM 000788 FEM1B NM 015322 FLJ14840 NM 032850 GCKR NM 001								FDX1	NM 004109	FLJ14642 NM 032818	GAPD	NM 002046
AKZ NM. 001625 BCS1L NM. 004328 CGF01 NM. 015935 DC8 NM. 015471 FEM1A NM. 020177 FLJ14827 NM. 032848 GCHFR NM. 005 AKAP13 NM. 007200 BET1 NM. 005868 CGF11 NM. 015941 DCK NM. 000788 FEM1B NM. 015322 FLJ14840 NM. 032850 GCKR NM. 001	ISG NM 001622	001622 BCL6	NM 001706	CGBP	NM_014593	DC13	NM 020188	FE65L2		FLJ14697 NM 032828		NM_000158 NM_005875
												NM_005258
AKR1C2 NM 001354 BF NM 001710 CGI-51 NM 015380 DCLRE1B NM 022836 FETUB NM 014375 FLJ20010 NM 019021 GDAP2 NM 017	R1C2 NM 001354	001354 BF	NM 001710	CGI-51	NM 015380	DCLRE1B	NM_022836	FETUB	NM, 014375	FLJ20010 NM_019021	GDAP2	NM_001486 NM_017686
AKR1C4 NM 001818 BIKE NM 017593 CHISLI NM 001278 DDA3 NM 032636 FHIT NM 002012 FL 20037 NM 017633 IGCCY NA 000	R1C4 NM 001818	001818 BIKE	NM 017593		NM_001278		NM 022487 NM 032636	FHIT	NM 000143 NM 002012	FLJ20014 NM 017622 FLJ20037 NM 017633		NM 005262 NM 000821
ALCAM NH 001627 BIRCS NM 016252 CHIC2 NM 012110 DDX18 NM 006773 FIGF NM 004469 FLJ20080 NM 017657 GIOT-2 NM 016							NM 006773	FIGF	NM 004469	FLJ20080 NM 017657	GIOT-2	NM 016234
ALDH2 NM 000690 BPHL NM 004332 CHP NM 007236 DDX28 NM 018380 FLJ10038 NM 017976 FLJ20084 NM 017659 GJA4 NM 002	DH2 NM 000690	000690 BPHL	NM 004332	CHP	NM 007236	DDX28	NM 018380	FLJ10038				NM_017655 NM_002060
ALUHJAN NM 000591 BRCA1 NM 007295 CIAO1 NM 004804 DDX35 NM 021931 FLJ10111 NM 017999 FLJ20123 NM 017674 GJB1 NM 000										FLJ20123 NM_017674	GJB1	NM_000166
ALDH5A1 NM 001080 BRIP1 NM 032043 CITED2 NM 005079 DDX8 NM 004941 FLJ10143 NM 018009 FLJ20130 NM 017681 GLYAT NM 005	DH5A1 NM 001080	001080 BRIP1	NM 032043	CITED2	NM 005079	DDX8	NM 004941	FW10143	NM 018009	FLJ20130 NM 017681	GLYAT	NM 020198 NM 005838
ALDOC NM 005165 BTF3 NM 001207 CKN1 NM 000082 DEDD2 NM 133328 FLJ10287 NM 019083 FLJ20287 NM 017746 GNB11 NM 053	DOC NM 005165	005165 BTF3	NM 001207	CKN1	NM 000082	DEDD2		FLJ10276		IFLJ20202 NM 017709 IFLJ20287 NM 017746		NM 001500 NM 053004
ALSZ NM 020919 BTG1 NM 001731 CKS2 NM 001827 DEPP NM 007021 FLJ10330 NM 018081 FLJ20331 NM 017768 GNG5 NM 005		020919 BTG1 057177 BTN2					NM 007021	FLJ10330	NM 018081	FLJ20331 NM 017768	GNG5	NM 005274
AMACR NM 014324 BYSL NM 004053 CLCN3 NM 001828 DJ37E16.5 NM 020315 FLJ10415 NM 018089 FLJ20452 NM 017828 GNS NM 002	MACR NM_014324	014324 BYSL	NM 004053	CLCN3	NM_001829	DJ37E16.5	NM 020315	FLJ10415	NM_018089	FLJ20452 NM 017828		NM 015960 NM_002076
AMBP NM, 001633 C12018 NM, 006817 CLCNB NM 001286 DJ726C3,2 NM, 025227 FLJ10422 NM, 018091 FLJ20511 NM, 017853 GOLGA2 NM, 004									NM 018091	FLJ20511 NM_017853	GOLGA2	NM_004488
AMT NM 000481 C140f3 NM 012111 CLON2 NM 020384 DKFZp434D177 NM 032264 FLJ10482 NM 018107 FLJ20580 NM 017887 GOLPH4 NM 014	T NM 000481	000481 C14or	13 NM 012111	CLON2	NM 020384	DKFZp434D177	NM 032264	FLJ10482	NM 018107	FLJ20580 NM 017887	GOLPH4	NM 002078 NM 014498
ANG NM 001145 C1018 NM 004872 CLDN3 NM 001306 DKFZP434H0115 NM 031421 FLJ10511 NM 018120 FLJ20595 NM 017894 GOSR2 NM 004 ANKRA2 NM 023039 C1S NM 001734 CLONE24922 NM 015679 DKFZP434J037 NM 030952 FLJ10525 NM 018126 FLJ20619 NM 017904 GOT1 NM 002	KRA2 NM 023039	023039 C1S	NM 001734								GOSR2	NM 004287 NM 002079
ANPEP NM 001150 C2 NM 000063 CLPTM1 NM 001294 DKFZP434L0117 NM 022778 FLJ10535 NM 018129 FLJ20627 NM 017909 GPC6 NM 005	PEP NM_001150	001150 C2	NM 000063	CLPTM1	NM 001294	OKFZP434L0117	NJJ 022778	FLJ10535	NM 018129	FLJ20627 NM 017909	GPC6	NM_005708
ANXA6 NM 001155 C2001154 NM 032485 CLTA NM 001833 DKFZP564G2022 NM 015497 FLJ10583 NM 018148 FLJ20671 NM 017924 GPR39 NM 001	XA6 NM 001155	001155 C20or	[154 NM 032485	CLTA	NM 001833	DKFZP564G2022	NM 015497	FLJ10583	NM_018146	FLJ20628 NM_017910 FLJ20671 NM 017924		NM_020806 NM_001508
ANXA9 NM 003568 C20ad163 NM 080749 CLTCL1 NM 001835 DKFZP564L2423 NM 030805 FLJ10604 NM 018154 FLJ20699 NM 017931 GPT NM 005						DKFZP564L2423	NM 030805	FLJ10604	NM 018154	FLJ20699 NM 017931	GPT	NM 005309
AP2A1 NM 130787 C20off172 NM 024918 CNOT2 NM 014515 DKFZP564O0523 NM 032120 FLJ10640 NM 019023 FLJ20718 NM 017939 GPX2 NM 002	2A1 NM 130787	130787 C20or	1172 NM, 024918	CNOT2	NM 014515	DKFZP56400523	NM 032120	FLJ10640	NM 019023	FLJ20718 NM_017939	GPX2	NM_000581 NM_002083
AP3B1 NM_003664 C200f1188 NM_015638 CNQT4 NM_013316 DKFZP566C243 NM_015388 FLJ10661 NM_018172 FLJ20729 NM_017953 GRHPR NM_012 AP3M1 NM_012095 C200f32 NM_020356 COASTER NM_015555 DKFZP566M1046 NM_032127 FLJ10761 NM_018208 FLJ20730 NM_017945 GRIK3 NM_000	3M1 NH 012095	012095 C20or	132 NM 020356			LDKFZP568C243	NM_01538B NM_032127				GRHPR	NM 012203 NM 000831
AP4B1 NM_006594 C200fl4 NM_015511 COPB NM_016451 DXFZp5660084 NM_015510 FLJ10774 NM_024662 FLJ21007 NM_030794 GRIN2D NM_0000	4B1 NM_006594	006594 C20or	4 NM_015511	COPB	NM, 016451	DXFZp5660084	NM 015510	FLJ10774	NM 024662	FLJ21007 NM 030794	GRIN2D	NM 000836
APCS NM. 001639 C20017 NM 024120 COPS78 NM_022730 DKFZP586A0522 NM_014033 FLJ10871 NM_018250 FLJ21272 NM_025032 GSK3B NM_002	CS NM 001639	001639 C20or	77 NM 024120	COPS7B		DKFZP586A0522						
APEH NM 001640 C20orf72 NM 052865 COQ3 NM 017421 DKFZP586J0119 NM 015636 FLJ10801 NM 018260 FLJ21415 NM 024738 GSPT1 NM 002			172 NM 052865	COO3	NM 017421	DKFZP586J0119	NM 015636	FLJ10801	NM 018260	FLJ21415 NM 024738	GSPT1	NM 002094
APG3 NM 022488 C210rf18 NM 017438 C0X7A2 NM 001865 DLEU1 NM 005087 FLJ11011 NM 018299 FLJ21908 NM 024604 GSTA4 NM 0018	G3 NM 022488	022488 C21or	118 NM 017438	COX7A2	NM, 001865	DLEU1	NM_005887	FLJ11011	NM 018299	FLJ21908 NM 024604	GSTA4	NM_000178 NM_001512
APOA1 NM 009039 C2F NM 006331 CPB2 NM 016413 DMC1 NM 007068 FLJ11046 NM 018709 FLJ21979 NM 022461 G5TT1528 NM 0447	OA1 NM 000039	000039 C2F	NM 006331	COX7A2L CPB2			NM 001933 NM 007068	JFLJ11029	NM 018304	FLJ21934 NM 024743	IGSTM4	NM 000850
I TRALE THE ALERA TAKE THE COLUMN	DA2 NM 001643	001643 C3	NM 000064	CPSF5	NM 007006		NM 005880	FLJ11159	NM_018343	FLJ21963 NM_024560	GTF2ET	NM_005513

Fig. 18B

Gana Name	RelSeq	·· Gene Nam	a RefSeq	Gene Nam	e. Reiseq	Gene Nam	ier RēfSēdu	Gens Nar	ne RaiSeo	T- Gene Na	me RafSeq . #D	Gene Nam	va Baltan Til
GTF2H1 GTP8G3	NM 005316 NM 032620	IGF1 IGFBP1	NM 000618 NM 000596	LOC51015 LOC51016	NM, U15U48	METAP2	NM 006838 NM 054013	MRPL49 MRPL51	NM_004927	OSGEP	NM_017807	PPP1R12	3 NM_032105
GYS2 H2A/S	NM 021957	IL11RA	NM_004512	LOC51028	NM 016072	MGC:1337	9 NM 016499	MRPS11	NM 016497 NM 022839	OSMR OTC	NM 003999 NM 000531	PPP1R158	NM 024607
H2AFG	NM 080596 NM 021065	IL15 IL1RAP	NM 000585 NM 002182	LOC51027 LOC51054	NM_016074 NM_015899	MGC10433 MGC10703	3 NM_024321 2 NM_032663	MRPS14	NM_022100 NM_016065	p100 P115	NM_014390 NM_003715	PPP1R3C PPP1R3D	NM_005398 NM_006242
H2AFO H2BFA	NM 003516 NM 003518	IL22R IL2RB	NM 021258 NM 000878	LOC51060 LOC51064	NM 015913 NM 015917	MGC10823 MGC10924	3 NM 031437	MRPS186	3 NM 014048	p21UAS1	1 NM 000389	PPP2CA	NM 002715
H2BFB H2BFF	NM ,021063	IL6ST	NN 002184	LOC51074	NM_015957	MGC10940	NM_032303	MRPS21	NM_018997	p21UAS8 p21UAS9	NM_000389	PPP2R5B PPP4R1	NM_006244 NM_005134
H2BFG	NM 021062 NM 003522	IMMT INADL	NM 006839 NM_005799	LOC51091 LOC51096	NM 016955 NM 016001	MGC10960 MGC10974		MRPS30 MRPS35	NM 016640 NM 021821	P23 P29	NM 006601 NM_015484	PPP5C	NM 006247 NM 005710
H326 H4F2	NM_015726 NM_003548	INHBC	NM_005538 NM, 014425	LOC51104 LOC51107	NM 016014 NM 016022	MGC10999 MGC11034	NM_032307	MRPS38	NM_033281	P2RY2	NM 002564	PRCC	NM_005973
H4FD H6PD	NM 003541	IRF6	NM 006147	LOC51134	NM 016122	MGC11268	NM 024322	MRPS7 MRS2L	NM_015971 NM_020662	PABPC1 PABPN1	NM_002568 NM_004643	PRCP PRKAB2	NM_005040 NM 005399
HAAO	NM 004285 NM 012205	ITGA6 ITGAL	NM 000210 NM_002209	LOC51142 LOC51143	NM 016139 NM 016141	MGC1127 MGC11279	NM 033549 NM_024326	MST1 MSTP028	NM 020998 NM 031954	PAFAH2 PAI-R8P1	NM 000437 NM 015640	PRKCABP PRKCL2	NM 012407 NM 006256
HADH2 HADHA	NM 004493 NM 000182	ITIH3 ITIH4	NM 002217 NM 002218	LOC51174 LOC51175	NM 016261 NM 016262	MGC12435 MGC12943	NM 031427	MT1H MT1L	NM 005951	PAK4	NM 005884	PRLR	NM 000949
HADHB HADHSC	NM 000183	ITM1	NM_002219	LOC51187	NM 016304	MGC12981	NM 032357	MT1X	NM_002450 NM_005952	PAUMD PANK	NM_017734 NM_138318	PRO1728 PRO2389	NM_018505 NM_025230
HAL	NM 005327 NM 002108	JIK	NM_002223 NM_016281	LOC51205 LOC51231	NM 016361 NM_016440	MGC13008 MGC13017	NM_032686 NM_080656	MT2A MTHFD1	NM 005953 NM 005956	PARVB PAX8	NM_013327 NM_013952	PRO2831	NM 018540 NM 003891
HAO1 HARC	NM, 017545 NM, 017913	JRKL JUN	NM_003772 NM_002228	LOC51240 LOC51246	NM 016467 NM 016479	MGC13033 MGC13102	NM 031447	MTHFR MTHFS	NM_005957	PBEF	NM_005746	PRPF31	NM_015829
HAX1 HBP1	NM 006118 NM 012257	JunB(-)1kb JunB(-)2kb	NM 002229 NM 002229	LOC51285	NM_016563	MGC13138	NM_033410	MTMR2	NM_006441 NM_003912	PC0H20 PCK1	NM 022843 NM 002591	PRPS1 PRSS25	NM 002764 NM 013247
HBQ1	NM_005331	JunB(-)3kb	NM,002229	LOC51287 LOC51292	NM 016565 NM 016576	MGC13159 MGC1346	NM_032758	MTMR4 MTP	NM 004687 NM 000253	PCK2 PCMT1	NM 004563 NM_005389	PSA PSMA1	NM 021154 NM 002786
HBS1L HBXIP	NM_006620 NM_006402	KAP3A KBRAS1	NG 000941 NM 020345	LOC51326 LOC51596	NM_016632 NM_015921	MGC14151 MGC14421	NM_032356 NM_032907	mtTFB MUT	NM_016020 NM_000255	PCYT1A PDCD4	NM,005017	PSMA2	NM_002787
HCA112 HCDI	NM 018487 NM 020195	KCNC3 KCNJ12	NM 004977 NM 021012	LOC51601 LOC51611	NM 015929	MGC14433	NM 032904	MYO1A	NM 005379	PDE11A	NM_014456 NM_016953	PSMA5 PSMD10	NM 002790 NM 002814
HDAC6	NM_006044	KCNN2	NM_021614	LOC51633	NM 015958 NM 016023	MGC14839 MGC14844	NM 032341	N8AMT1 NAGA	NM_013240 NM_000262	PDE4DIP PDE6D	NM_014644 NM_002601	PSMD7 PSME3	NM_002811 NM_005789
HDAC6 HEL308	NM 006044 NM 133636	KEO4 KHORBS1	NM 006459 NM 006559	LOC51644 LOC51651	NM 016057 NM_016077	MGC15435 MGC15504	NM 032367 NM 032751	NAGK NAPA	NM 017567 NM 003827	PDIR PDK2	NM 006810 NM 002611	PT0012 PT0013	NM 014039 NM 015952
HEXA HEY1	NM_000520 NM_012258	KIAA0092 KIAA0102	NM 014679 NM 014752	LOC51659 LOC54516	NM 016095 NM 019041		NM 138570	NAT8 NBP	NM 003960	PDK4	NM_002612	PT0015	NM_014040
HFL3 HGC6.2	NM 005666	KIAA0103	NM 014673	LOC54518	NM 019043	MGC15677	NM 032878	NCALD	NM_025233 NM_032041	PDZK1 PECI	NM .002614 NM .006117	PTK2 PTPN18	NM_005607 NM_014369
HGD	NM_014356 NM_000187	KIAA0105 KIAA0141	NM 004908 NM 014773	LOC55580 LOC55815	NM_017571 NM_018430	MGC15737 MGC15906		NCBP1 NCBP2	NM_002488 NM_007362	PELO PEMT	NM_015946 NM_007169	PTPN4 PTPRE	NM_002830 NM_006504
HIF1A HINT2	NM_001530 NM_032593	KIAA0205 KIAA0255	NM 014873 NM 014742	LOC55954 LOC56834	NM 019103 NM 020155	MGC16733 MGC16943	NM_033547	NCF1 NCK1	NM 000265 NM 006153	PEPD	NM, 000285	PTPRG	NM_002841
HKE2 HKE4	NM_014260 NM_006979	KIAA0258 KIAA0266	NM 014785 NM 021645	LOC56902 LOC57018	NM 020143	MGC17347	NM 138333	NCOA3	NM 006534	PEX11B PEX13	NM 003846 NM 002618	PURG PWP1	NM 013357 NM_007062
HLA-B	NM 005514	KIAA0391	NM 014672	LOC57019	NM_020307 NM_020313	MGC19595 MGC2404	NM 032360	NCOA5 NCOR1	NM 020967 NM 006311	PEX16 PEX3	NM_057174 NM_003630	PYGL PZP	NM 002863 NM 002864
HLA-F HMCS	NM_018950 NM_017947	KIAA0409 KIAA0433	NM_015324 NM_015216	LOC57107 LOC57228	NM_020381 NM_020467	MGC2474 MGC2477	NM 023931 NM 024099	NDRG1 NDUFA4	NM_006096 NM_002489	PFKFB4 PGM1	NM 004567 NM 002633	QP-C R3HDM	NM 014402 NM 015361
HMG1 HMG17L3	NM_002128 NM_006353	KIAA0438 KIAA0618	NM_014819 NM_014833	LOC57406 LOC57828	NM_020676 NM_021183	MGC2488 MGC2560	NM 024039 NM 031452	NDUFA6	NM_002490	PHACS	NM_032592	RA410	NM, 016106
HMOX2	NM 002134	KIAA0645	NM 014662	LOC57862	NM_021188	MGC2629	NM 032522	NDUF81 NDUF85	NM 004545 NM_002492	PHLDA1 PHTF1	NM_007350 NM_006608	RAB10 RAB11A	NM_016131 NM_004653
HNF4a7 HNMT	AF509467 NM_006895	KIAA0660 KIAA0670	NM 012297 NM 014977	LOC64182 LOC81034	NM 022359 NM 030780	MGC2650 MGC2734	NM 024108 NM 033117	NDUFS1 NDUFS4	NM 005008 NM_002495	PIGPC1 PIGPC1	NM 022121 NM, 022121	RAB18 RAB2	NM 021252 NM 002865
HNRPA1 HNRPR	NM 031157 NM 005826	KIAA0747 KIAA0792	NM 015292 NM 014698	LOC81558 LOC84518	NM 030802 NM 032488	MGC2747 MGC2835	NM 024104 NM 024072	NEDD8 NEK2	NM 006156 NM_002497	PIGPC1	NM 022121 NM 022121	RAB30	NM 014488
HOOK3 HOXA1	NM_032410 NM_005522	KIAA0795 KIAA0806	NM 025010 NM 014813	LOC84661 LOC89953	NM_032574 NM_138343	MGC3067	NM 024295	NET-2	NM_012338	PIGS	NM 033198	RAB33B RAB4B	NM_031296 NM_016154
HOXC8	NM 022658	KIAA0872	NM 014940	LOC90799	NM 138363	MGC3180 MGC3222	NM 024041 NM 024334	NFE2L1 NFKBIB	NM 003204 NM 002503	PIK3R3 PIK4CB	NM_003629 NM_002651	RAB6KIFL RAB9P40	NM_005733 NM_005833
HPCL2	NM 012260 NM 002151	KIAA0905 KIAA0914	NM 014933 · NM 014883	LOC91689 LR8	NM 033318 NM 014020	MGC3248 MGC3413	NM_032486 NM_032678	NFKBIB NFKBIB	NM_002503 NM_002503	PILB PINK1	NM_012228 NM_032409	RABEX5 RAD17	NM 014504 NM 133338
HPRP4P HPX	NM 004697 NM 000613	KIAA1017 KIAA1041	NM 007216 NM 014947	LSM3 LSR7	NM 014463 NM_018559	MGC4161 MGC4189	NM 024303 NM 032308	NFKBIB NFYA	NM 002503 NM 002505	PIP5K1A PIPOX	NM 003557	RAD238	NM 002874
HRIHFB2436 HSA011916		KIAA1116 KIAA1169	NM 014892 NM 017901	LTA4H	NM_000895	MGC4400	NM_032679	NKTR	NM_005385	PIR	NM, 016518 NM_003662	RAD50 RAGA	NM 133482 NM 006570
HSD11B1	NM 005525	KIAA1453	NM 025090	LZTR1 M17S2	NM 006767 NM 031858	MGC4606 MGC4638	NM 024516 NM 031479	NME1 NOLC1	NM_000269 NM_004741	PIST	NA1_020399 NM 012399	RA-GEF-2 RAMP	NM_016340 NM_016448
HSD1782 HSD1784	NM_002153 NM_000414	KIAA1638 KIF1B	NM 025132 NM 015074	M96 MADCAM1	NM 007358 NM 007164	MGC4663 MGC4677	NM_024514 NM_052871	NONO NPAS2	NM_007363 NM_002518	PKM2 PLA2G13	NM_002654 NM_032562	RANBP8 RANGAP1	NM_008390 NM_002883
HSD17B7 HSPA5	NM 016371 NM 005347	KIF9 KLF15	NM 022342 NM 014079	MADH4 MAF	NM 005359 NM 005360	MGC4767 MGC5302	NM 032314 NM 024089	NPAT NPC1	NM 002519 NM 000271	PLAB PLAGL2	NM 004864	RAP1GA1	NM 002885
HSPC002 HSPC048	NM 015362 NM_014148	KLHL6 KNG	NM, 130446	MAGOH	NM_002370	MGC5509	NM_024093	NROB2	NM_021969	PLD2	NM_002657 NM_002663	RASA1 RASSF1	NM 022850 NM 007182
HSPC051	NM 013387	KNSL4	NM 000893 NM 007317	MAL2 MANBA	NM_052886 NM_005908	MGC9084 MGEA5	NM_033418 NM_012215	NR1H3 NR1I2	NM_005693 NM_022002	PLGL PLSCR1	NM_002665 NM_021105	RBBP4 RBM15	NM 005810 NM 022768
HSPC052 HSPC111	NM 014150 NM 016391	KPNB1 KRT10	NM 002265 NM 000421	MAOA MAP3K11	NM 000240 NM 002419	MGST1 MGST2	NM_020300 NM_002413	NR3C1 NR5A2	NM 000176 NM 003822	PME-1 PMS1	NM_016147 NM_000534	RBM6 RBM7	NM 005777
HSPC117 HSPC129	NM_014306 NM_016396	LAD1 LALP1	NM_005558 NM_020354	MAP3K4 MAP3K7	NM 005922 -	MGST3	NM 004528	NRAS	NM_002524	PMS2	NM, 000535	RBP5	NM 016090 NM_031491
HSPC141	NM 014172	LAPTM4A	NM 014713	MAP7	NM 003188 NM 003980	MIPEP MLC1SA	NM 005932 NM 002475	NRCAM NRD1	NM 005010 NM 002525	PMS2L8 PNAS-131	NM 005394 NM 031446	RB\$K RBT1	NM_022128 NM_013368
HSPC154 HSPC157	NM_014177 NM_014179	LATS1 LBP	NM 004690 NM 004139	MAPK7 MAT1A	NM_002749 NM_000429	MNAT1 MOV10	NM_002431 NM_020963	NS1-BP NT5C3	NM_006469 NM_016489	PNKP PNLIPRP1	NM, 007254	RCL RDBP	NM_006443 NM_002904
HSPC166 HSPC213	NM 014186 NM 016475	LC27 LCN2	NM 018407 NM 005564	MAT2A MBD4	NM 005911 NM 003925	MPP1 MPPE1	NM 002436 NM 023075	NTHL1 NTN4	NM 002528 NM 021229	POLB	NM 002690	RDH5	NM 002905
HSU79274 HSU84971	NM 013300 NM 013303	LENG5	NM_024075	MCCC1	NM_020166	MRE11A	NM 005590	NUDT2	NM 001161	POLD4 POLE3		REA RECOLS	NM_007273 NM_004259
HT002	NM 014066	LEPR LGALS1	NM, 002303 NM 002305	MCEE MCP	NM 032601 NM 002389	MRP63 MRPL15	NM 024026 NM 014175	NUDT5 NUFIP1	NM 014142 NM 012345	POLR2A POLR2K		RENT1 RFC3	NM 002911 NM 002915
HT007 HT010	NM 018480 NM_018471	LIMK2 LISCH7	NM 005569 NM 015925	MDFI MDH1	NM 005586 NM_005917	MRPL18 MRPL2	NM 014161 NM 015950	NUP107 NUP62	NM 020401	POLS PON1	NM 006999	RFC5 RGL	NM 007370
HT012 humNRDR		LIV-1 LNPEP	NM 012319 NM 005575	MDM2UAS6 MDM2UAS8	NM 002392	MRPL24	NM 024540	NUP98	NM 005387	POP5	NM 015918	RIG-I	NM 015149 NM 014314 NM 013400
HYAL3	NM 003549	LOC115330	NM_138445	MDS009	NM 020234	MRPL33 MRPL34	NM_004891 NM_023937	OAS1 OAS3	NM_002534 NM_006187	PORIMIN POV1	NA1_003627	RNASE2	NM_013400 NM_002934
IER5 IFITM2	NM 016545 NM 006435	LOC129401 LOC151534	NM 138482	MDS025 MDS029	NM 021825 NM 018464	MRPL37 MRPL4	NM_016491 NM_015958	OAZ2 OPA3	NM 002537	PP5395 PPFIBP1	NM_021732	RNASE3 RNASE4	NM_002935 NM_002937
IFNAR1 IFNGR1	NM 000629	LOC151636	NM 138287	MEA MEF2B	NM_014823 NM_005919	MRPL44 MRPL46	NM 022915 NM 022163	ORC3L ORM1	NM_012381	PPGB	NM_000308	RNF29	NM 033058
IFRD1	NM 001550	LOC51011	NM 016044	MEP50	NM 024102	MRPL48	NM 016055	ORM2	NM 000607	PPM1D PPP1R11	NM 003620 NM 021959	RNF5 RNGTT	NM 006913 NM 003800

Fig. 18C

Gene Name RNPC2	NM_004902	SLC25A13	NM_014251	TDRKH	NM 006862	Gene Name	NM 007259
RNPEPL1	NM 018226	SLC25A18	NM 031481	TEAD3	NM 00862 NM 003214	VPS45A	NM 000638
ROCKI	NM_005406	SLC25A5	NM 001152	TED	NM_015686	WASF3	NM_006646
RORC	NM_005060	SLC26A1	NM 022042	TEF	NM_003216	WASL	NM 003941
PC32	NM 006467	SLC2A8	NM_014580	TEGT	NM_003217	WBP4	NM 007187
PL18	NM 000979	SLC31A1	NM 001859	TESK2	NM 007170	WDF2	NM 052950
PL31	NM 000993	SLC35A2	NM 005660	TF	NM 001063	WDR10	NM 052985
PL37AP1	NG 000988	SLC35A3	NM 012243	THPO	NM_000460	WDR12	NM_018256
PL5	NM_000969	SLC38A1	NM, 030674	THTP	NM_024328	WDR13	NM_017883
PL7	NM_000971	SLC38A4	NM 018018	TIA1	NM_022037	XDH	NM 000379
IPLP1	NM 001003	SLC39A1	NM_014437	TIMM17A	NM_006335	XPA	NM_000380
PS16	NM_001020	SLC5A3	NM_006933	TIMM17B	NM_005834	XPC	NM 004628
PS19	NM_001022	SLC7A2	NM_003046	TIMM23	NM 006327	XPR1	NM 004736
IPS27A	NM 002954	SLC7A9	NM 014270	TIMM9	NM 012460	XRCC5	NM 021141
UPS3A	NM 001006	SLPI	NM_003064	TLH29	NM_032036	YKT6	NM_006555
RPS6KA5	NM_004755	SMAC	NM .019887	TLN1	NM_006289	YWHAB	NM_003404
PS6KB1	NM 003161	SMAP	NM_006695	TM4SF4	NM_004617	ZAN	NM_003386
QCD1	NM_005444	SMARCA5	NM_003601	TM9SF2	NM_004800	ZBRK1	NM_021632
ISHL1	NM_030785	SMARCE1	NM 003079	TMEM7	NM, 031440	ZF5128	NM 014347
ISP3	NM 031924	SMC2L1	NM 006444	TMF1	NM 007114	ZFP95	NM 014569
SU1	NM 012425	SMPD1	NM_000543	TMOD2	NM_014548	ZK1	NM_005815
TCD1	NM.003729	SNAI2	NM_003068	TMP21	NM, 006827	ZNF133	NM, 003434
TP801	NM_019058	SNAP23	NM 003825	TNFAJP1	NM, 021137	ZNF144	NM, 007144
UVBL2	NM_006666	SNAPC1	NM_003082	TNFRSF118		ZNF146	NM_007145
XRB	NM 021976	SNK	NM 006622	TNFRSF6	NM_000043	ZNF147	NM 005082
100A9	NM 002965	SNRPA	NM_004596	TNFRSF6	NM 000043	ZNF155	NM 003445
AA1	NM_000331	SNRPD3	NM_004175	TNFRSF6	NM 000043	ZNF183	NM_006978
	- NM_000331	SNRPF	NM, 003095	TNFRSF6	NM_000043	ZNF192	NM_006298
AA1	NM_000331	SWW1	NM_012245	TNFSF13	NM_003808	ZNF207	NM 003457
AA1	NM 000331	SNX1	NM 003099	TNS	NM 022648	ZNF214	NM 013249
AA2	NM 030754	SNX17	NM 014748	TOM1	NM 005488	ZNF22	NM 006963
AC	NM_018417	SNX3	NM 003795	TOMM70A	NM,014820	ZNF221	NM_013359
AD1	NM_006590	SNX5	NM_014426	TP53TG1	NM_007233	ZNF222	NM_013360
AMHD1	NM_015474	SOD1	NM, 000454	TPP2	NM_003291	ZNF224	NM 013398
AP18	NM 005870	SORCS3	NM 014978	TPT	NM 014317	ZNF225	NM 013362
AS10	NM_020368	SOX10	NM ,006941	TRA1	NM_003299	ZNF226	NM 016444
CAMOL	NM 006745	SP2	NM, 138406	TRAF6	NM_004620	ZNF230	NM_006300
CA2 CAND1	NM 002973	SPATA2	NM_006038	TRAP150	NM 005119	ZNF237	NM_014242
CANDI	NM 033630	SPATA6	NM 019073	TRIM15	NM 033229	ZNF281	NM 012482
CYA14	NM 005063 NM 032962	SPC18 SPOCK	NM 014300	TRIM26	NM 003449	ZNF302	NM 018443
CYA15	NM_032964	SPP2	NM 004598	TRIM31	NM 052816	ZNF361	NM_018555
CYA16	NM_004590	SQRDL	NM 006944 NM_021199	TRIM34 TRIM4	NM_130389	ZNF9	NM_003418
CYE1	NM_004757	SREBF2			NM 033017		NM_014415
DC1	NM_002997	SRP54	NM_004599	TRIP11 TRN-SR	NM_004239	ZNRD1	NM_014596
DCCAG10	NM_005869	SRP68	NM 003136		NM_012470	ZnTL2	INM. 133496
DCCAG18	NM_005645	SRPR	NM_014230	TRPC5	NM_012471		
DFR1	NM 012428	SSA2	NM, 003139	TRPS1 TSG101	NM_014112	1	
EC10L1	NM 006544	SSAT2	NM 004600 NM_133491	TSLRP	NM 006292		
EC23A	NM_006364	SSSCA1	NM_006396	TTY14	NM 012472 NM 031932		
EC24D	NM_014822	SSTR1	NM_001049	TUBB5	NM_006087		
EC61B	NM_006808	STAF42	NM_053053	TUFT1	NM_020127	1	
EL1L	NM 005065	STAF65(gamma)	NM 014860	TXNIP	NM 006472	Ì	
EMA3C	NM_006379	STAM	NM, 003473	TXNL	NM_004786		
EMA6C	NM_030913	STAM2	NM 005843	TXNRD1	NM 003330		
EMA7A	NM 003612	STARD7	NM 020151	TYMS	NM 001071	İ	
ENP1	NM_014554	STATI	NM_007315	U2AF1	NM ,006758	1	
EPX1	NM 016332	STAU2	NM 014393	U3-55K	NM 004704		
ERPINA1	NM_000295	STCH	NM 006948	U5-116KD	NM_004247		
ERPINA10	NM 016186	STIMI	NM_003156	UBE2B	NM_003337		
RPINA5	NM_000624	STK19	NM_004197	UBE2D3	NM_003340		
ERPINA6	NM_001758	STK2	NM_003157	ŬBE2M	NM 003969		
ERPINC1	NM_000488	STOML1	NM_004809	UBP1	NM_014517		
ERPIND1	NM 000185	STRAIT11499	NM 021242	UBOLN1	NM 053067		
ERPINE1	NM 000602	STX18	NM 016930	UBQLN2	NM_013444	1	
RPING1	NM_000062	SUCLA2	NM, 003850	UCH37	NM 015984		
RPINI1	NM_005025	SUCLG1	NM_003849	UCHL3	NM 006002		
S2	NM_031459	SUDD	NM_003831	UGDH	NM 003359	1	
3A3	NM_006802	SULT1A1	NM_001055	UGT2B11	NM 001073	1	
3B2	NM ,006842	SULT2A1	NM 003167	UGT2B15	NM.001076	1	
RS11	NM 004768	SUOX	NM 000456	UGTREL1	NM 005827		
RS5	NM 006925	SUPT3H	NM_003599	UGTREL7	NM 015139	ł	
RS8	NM 004592	SUPT5H	NM 003169	ULBP3	NM 024518		
3K	NM 005627	SUPV3L1	NM_003171	UPB1	NM 016327	1	
SK2	NM 016276	SYN3	NM 133632	UQCRC2	NM_003366	1	
ST1	NM 006704	SYTL4	NM 080737	URKL1	NM 017859	1	
12D3C	NM_005489	SZF1	980310, MM	UROD	NM 000374	1	
H3BGRL2	NM, 031469	TADA3L	NM, 133480	UROS	NM.000375	1	
LV '	NM 006928	TAF2GL	NG 001012	USP1	NM 003368	1	
X2	NM_016932	TAGLN2	NM_003564	USP15	NM_008313	1	
(B1	NM 006109	TARS	NM 003191	USP2	NM 004205	i	
(D1	NM, 004869	TAT	NM 000353	UXT	NM 004182	ĺ	
(RP1	NM_080876	TCF1	NM 000545	VAMP1	NM 014231	İ	
.C10A1	NM 003049	TCF12	NM 003205	VAMP5	NM_006634		
.C17A2	NM, 005835	TCF19	NM_007109	VDAC1	NM_003374		
.C17A5	NM 012434	TCF21	NM 003206	VDAC2	NM 003375		
	NM 025243	TCF7L2	NM 030756	VEGFC	NM 005429		
.C19A3			1411 000100				
.C19A3 .C22A1LS	NM_007105	TCIRG1	NM 006019	VEZATIN	NM_017599	•	
.C19A3							

(29/41)

Fig. 19A

	e RefSeq - 3		RefSeq	Gene Name	ReiSeq	Gene Name	RefSed	Gene Nam	e RelSer	Gene Nam ReiSeo	Gene Name Raiseq
101F6 4E-T	NM_007022 NM_019843	BIG1 BLTR2	NM 008121 NM 019839	CGBP CGI-01	NM 014593 NM 015935	DKFZP547N043 DKFZP584G2022	NM 032018	FW10477	NM_018105	FW20420 NM 017812	GPRK2L NM_005307
AAMP	NM .001087	BLZF1	NM_003665	CGI-203	NM 020408	DKFZP56410422	NM, 015497 NM, 031435	FLJ10509 FLJ10511	NM 018119 NM 018120	FLJ20422 NM 017814 FLJ20450 NM 017827	GRIK3 NM 000831 GRTH NM 013264
ABCB10 ABCB8	NM_012089 NM_007188	8M-002 8MH	NM 016617 NM 005180	CHEK2	NM 015380 NM 007194	DKFZP564L2423 DKFZP564M082	NM 030805 NM 014042	FLJ10525 FLJ10535	NM_018126	FLJ20498 NN1_019040	GRWD NM 031485
ABCB9	NM_019624	BMP5	NM,021073	CHERP	NM 008387	DKFZP56400463	NM 014156	FLJ10581	NM 018129	FLJ20508 NM 017850 FLJ20511 NM_017853	GSPT1 NM 002094 GSS NM 000178
ABCC5 ABCG1	NM 005688 NM 004915	BNC BNIP1	NM_001717 NM_001205	CHIC2 CHM	NM 012110 NM 000390	OKFZP58400523 DKFZP566B183	NM_032120 NM 015509	FLJ10583	1018148	FLJ20546 NM_017872	GSTZ1 NM 001513
ABH	NM_006020	BPGM	NM ,001724	CHMP1.5	NM_020412	DKFZP586C243	NM_015388	FLJ10628	NM 018154 NM 018159	FLJ20558 NM 017880 FLJ20624 NM 017906	GTF28 NM 001514 GTF2E1 NM 005513
ABS ABT1	NM_016222 NM_013375	BRAP BRCA1	NM 008768 NM 007295	CHRNB2 CIAO1	NM 000748 NM 004804	DKFZP566D1346 DKFZP566E144	NM 030816 NM 015523	FLJ10634 FLJ10637	NM_018163 NM_018164	FLJ20627 NM 017909 FLJ20628 NM_017910	GTF2H1 NM_005316
ACAD8	NM_014384	BRF2 BRIX	NM .018310	CIP29	NM_032364	DKFZP586A011	NM, 015416	FLJ10640	NEA_019023	FLJ20643 NM_017916	GTF2H4 NM_001517
ACADSB ACATN	NM 001609 NM 004733	BST1	NM 018321 NM_004334	CIR CITED2	NM 004882 NM 006079	DKFZP586J0119 DKFZP781E2110	NM 015636 NM_030953	FLJ10661 FLJ10774	NM 018172 NM 024662	FLJ20644 NM 017817 FLJ20651 NM 017919	GTF2I NM 033003 GTF3C5 NM 012087
ACO2 ACOX1	NM_001098 NM_004035	BTD BTRC	NM_000060 NM_033637	CKAP1 CKS2	NM_001281	DKFZp761J139 DKFZP7021166	NM_032280	FLJ10803	NM_018224	FLJ20671 NM 017924	GUSB NM. 000181
ACOX3	NM 003501	BUBIB	NM_001211	CLLD8	NM 001827 NM 031915	DLG4	NM 020441 NM 001365	FLJ10828 FLJ10853	NM 018233 NM 018246	FLJ20695 NM 017929 FLJ20729 NM 017953	H GS165L1 NM 004904 H17 NM 017547
IACP2 IACTR1A	NM 001610 NM_005736	BUB3 BYSL	NM_004725 NM_004053	CLONE24922 CLPTM1	NM 015679 NM 001294	DMAP1 DMP1	NM_019100	FLJ10856	NM D18247	FLJ20730 NM_017945	H326 NM 015726
AD-017	NM 018446	C11orf10	NM 014208	CLPX	NM 006660	DNAJB11	NM_004407 NM_016306	FLJ10871 FLJ10891	NM 018250 NM 018200	FLJ20731 NM 017945 FLJ20748 NM 019020	H3FM NM 021059 H4F2 NM 003548
AD022 AD034	NM 016614 NM 031480	C11orf2 C14orf3	NM 013265 NM 012111	CLTA CLTCL1	NM 001833 NM_001835	DNAJB12 DNAJB4	NM 017626 NM 007034	FLJ10989 FLJ10998	NM 018292 NM_018294	FLJ20772 NM 017956 FLJ20859 NM 022734	H4FI NM 003544
AD158	NM_032270	C1D	NM 006333	CNAP1	NM 014865	DPAGT1	NM 001382	FLJ11000	NM, 018295	FLJ21272 NM_025032	HASJ4442 NM 017528
AD24 ADAT1	NM 022451 NM 012091	C1ort22 C1ort25	NM 025191 NM 030934	CNOT3 CNOT4	NM 014516 NM 013316	DPH2L2 DPM1	NM 001384 NM 003859	FLJ11016 FLJ11017	NM 018301 NM 018302	FLJ21613 NM 021929 FLJ21742 NM 032207	HAX1 NM 006118 HBOA NM 007067
ADCY7	NIJ ,001114	C1 ori8	NM_004872	COASTER	NM_015555	DPM2	NM_003863	FLJ11029	NM_018304	FLJ21820 NM 021925	HBP1 NM 012257
ADD2 ADSS	NM 001617 NM 001126	C20orf1 C20orf10	NM_012112 NM_014477	COPB	NM_006710 NM_016451	DSCR3 DSCR5	NM 006052 NM 016430	FLJ11046 FLJ11159	NM_018309 NM_018343	FLJ21934 NM 024743 FLJ21939 NM 022461	HBQ1 NM 005331 HBXIP NM 006402
AF093680 AF140225	NM 013242 NM 030789	C20orf111 C20orf12	NM 016470 NM 018152	COPB2 COPS7A	NM_004766	DSS1	NM 006304	FUH1186	NM_018353	[FLJ21945 NM_025203	HCAP-G NM_022346
AF15Q14	NT# 020380	C20orf13	NM 017714	COPS78	NM 016319 NM 022730	DYRK1B E2F4	NM 004714 NM 001950	FLJ11193 FLJ11220	NM 018356 NM 018364	FLJ21952 NM 022484 FLJ21977 NM 032213	HCDI NM 020195 HCNGP NM 013260
AGA AGTPBP1	NIA 000027 NIA 015239	C20orf14 C20orf154	NM 012469 NM 032485	COX7A2 COX7A2L	NM 001865 NM 004718	E2F5 E2IG3	NM 001951 NM 014366	FLJ11271	NM 018373	FLJ21986 NM 024913	HD NM 002111
ALP	NM 003977	C20orf164	NM_080752	COX7C	NM_001867	EAF1	E80EE0, MM	FU11274 FLJ11292	NM_018375 NM_018382	FLJ22028 NM 024854 FLJ22169 NM 024085	HDAC8 NM 018485 HEC NM 006101
AK2 AKR1B1	NM 001625 NM 001628	C20orf188 C20orf28	NM 015638 NM 015417	COXB CPA2	NM_004074 NM_001869	EED EEF182	NM 003797 NM 021121	FLH1301 FLH1838	NM 018385 NM 024664	FLJ22184 NM_025094	HEL308 NM 133636
ALS2	NM 020919	C20orf30	NFA 014145	CPSF5	NM 007006	[EFG1	NM_024996	FLJ11848	NM 025155	FLJ22347 NM 022830	HEXA NM 000520 HGD NM_000187
AMSH ANKRA2	NM 006463 NM 023039	C20orf33 C20orf4	NM 030877 NM 015511	CPT1B CREBL1	NM 004377 NM 004381	EGLN2 EHD3	NM 053046 NM 014600	FLJ12085 FLJ12168	NM 022771 NM 024682	FLJ22501 NM 024747 FLJ22551 NM 024708	HHEX NM 002729 HHLA2 NM 007072
APIM1 AP2A1	NM 032493 NM 130787	C20orf43 C20orf44	NM,016407 NM 018244	CREBL2 CRFG	NM_001310	EIF1A	NM 001412	FLJ12455	NM_022078	FLJ22555 NM_024520	HIF1AN NM 017902
AP2B1	NM, 001282	C20orf45	NM_016045		NM_012341 NM_016507	EIF281 EIF2S1		FLJ12525 FLJ12571	NM 031206 NM 024926	FLJ22637 NM 025165 FLJ22688 NM 025129	HIRIP3 NM 003609
AP2M1 AP2S1	NM 004068 NM 021575	C20orf64 C20orf72	NM 033550 NM 052865		NM 004830 NM 001889	EIF2S2 (EIF2S3	NM_003908	FLJ12707 FLJ12735	NM_022067 NM_024857	FLJ22729 NM 024683	HKE4 NM_006979
AP3M1	NM_012095	C20or777	NM 021215	CRYZL1	NM 005111	EIF3S2	NM ,003757 .	FLJ12770	NM_032174	FLJ22865 NM 025109 FLJ22875 NM 032231	HLF NM 002126 HMG1 NM 002128
AP4B1 APACD	NM_006594 NM_005783	C21orf18 C21orf55	NM_017438 NM_017833		NM_004077 NM_004383	EIF3S6 EIF4G1		FLJ12765 FLJ12788	NM_024855 NM_022492	FLJ23109 NM_024814 FLJ23182 NM 022366	HMG2 NM 002129 HNRPA0 NM 006805
APC10 APG3	NM 014885 NM 022488	C21orf59 C2F	NM 021254 NM 006331	CSNK2A1	NM 001895	EIF5	NM_001969	FLJ12879	NM 024757	FLJ23251 NM 024818	HNRPA1 NM_031157
APMCF1	NM_021203	C2orf9	NM 032309		NM 001324 NM 015235	ELL EPHA1		FLJ12888 FLJ12895	NM 024945 NM 023926	FLJ23263 NM 025115 FLJ23468 NM 024629	HNRPC NM_031314 HPCL2 NM_012260
AQP3 AQP6	NM 004925 NM 001652	C3ori4 C4ori1	NM 019895 NM 006345		NM 001326 NM 001327	ERCC5 EWSR1	NM 000123	FLJ12910	NM 024573	FLJ23469 NM 024710	HPRP4P NM 004697
ARD1	NM 003491	C5orf6	NM 016605	CTMP	NM 053055	EXO1		FLJ12960 FLJ13102	NM 024638 NM_024887	FLJ23499 NM 022761 FLNA NM 001458	HRB2 NM 007043 HRMT1L2 NM 001536
ARF1GAP ARFD1	NM 018209 NM 001656	C6orf11 C6orf35	NM 005452 NM 018452		NM,001903 NM 003591	EZFIT F12	NM_021216	FLJ13158 FLJ13194	NM 024909 NM 025146	FNTB NM_002028	HSGT1 NM_007265
ARHGAP11	I. NM .014783	C7orl10	NM_024728	CXorf12	NM_003492	F23149, 1	NM 019088	FLJ13195	NM_022906	FOXO1A NM 002015	HSP105B NM 006644 HSPA5 NM 005347
ARL1 ARS2	NM_001177 NM_015908	C9orf12 C9orf5	NM_022755 NM_032012		NM_030579 NM_015247	FACTP140 FANCE		FLJ13220 FLJ13273	NM 021927 NM 024751	FRG1 NM 004477 FRSB NM 005687	HSPC003 NM 014017 HSPC016 NM 015933
ARSDR1 ASB3	NM 016026 NM 016115	CAP CAPZA2	NM 006367 NM 006136	CYP51	NM 000786	FBXO24	NM 012172	FLJ13291	NM 032178	FTL NM 000146	HSPC031 NM 016101
ASE-1	NM 012099	CAT56	NM 025263	D13S106E	NM 006023 NM 005800	FBXW2		FLJ13315 FLJ13491	NM_025005 NM_024623	FTSJ1 NM_012280 FUBP1 NM_003902	HSPC051 NM_013387 HSPC052 NM_014150
ATF4 ATF6	NM 001675 NM 007348	CAV1 CBARA1	NM 001753 NM 006077		NM_007158 NM_053281	FDPS FDX1		FLJ13611 FLJ13615	NM_024941 NM_025114	FXC1 NM 012192 FYCO1 NM 024513	HSPC056 NM, 014154
ATF7	NM_006856	CBX5	NM_012117	DAD1	NM_001344	FDXR	NM, 024417	FLJ13798	NM_024773	G10 NM 003910	HSPC072 NM 014162 HSPC111 NM 016391
ATP10C ATP5B	NN1_024490 NM_001686	CCNE1 CCNT1	NM_001238 NM_001240			FE65L2 FEN1	NM_006051 NM_004111	FLJ13912 FLJ13949	NM_022770 NM_025077	G22P1 NM 001469 G6PD NM 000402	HSPC117 NM 014306
ATP5F1 ATP5G3	NM 001688 NM 001689	CC168 CC17	NM 006584 NM 006429	DC13	NM 020188 NM 031210	FGF13	NM 004114	FLJ13962	NM 024862	GABPA NM 002040	HSPC129 NM 016396
ATP5J2	NM 004889	CCT8	NM_006585	DC8	NM_015471	fH	NM 002009 NM 000143	FLJ14431 FLJ14451	NM 032783 NM 032786	GABPB2 NM_002041 GABRE NM_021984	HSPC134 NM 014169 HSPC138 NM 016401
atp6e atp6m	NK1 001698 NM 015994	CDC10 CDC23	NM_001788 NM_004661	DCLRE1B	NM_022836	FHIT	NM 002012	FLJ14488 FLJ14511	NM_032792	GALNAC41 NM 031422	HSPC141 NM_014172
ATP6S14	NM 004231	CDC25A	NM 001789	DDOST	NM 005216	FKBP3	NM 002013	FLJ14547	NM 033087 NM 032804	GAS1 NM 002048 GBF1 NM 004183	HSPC142 NM 014173 HSPC144 NM 014174
AUP1 AUTL1	NM, 012103 NM, 032852	COC428P8 COC45L	NM_008035 NM_003504	DDX10 DDX21	NM 004398 NM 004728			FLJ14697 FLJ14803	NM_032826 NM_032842	GCN5L1 NM_001487 GDAP2 NM_017686	HSPC148 NM 016403 HSPC152 NM 016404
B3GNT6 BAD	NIA 005876 NIA 004322	CDC5L CDC5	NM 001253	DDX28	NM 018380	FLJ10038	NM 017978	FLJ14840	NM 032850	GHITM NM 014394	HSPC157 NM 014179
BAG4	NIJ 004874	CDCA1	NM 001254 NM 031423	DDXB	NM_004941	FLJ10116	NM 018000	FLJ14855 FLJ20010	NM 033210 NM 019021	GIOT-3 NM 016265 GJA4 NM 002060	HSPC160 NM 014182 HSPC166 NM 014186
BARD1 Bati	NM 000465 NM 004640	CDIPT COK5	NM 008319 NM 004935	DED		FLJ10142	NM_018008	FLJ20045 FLJ20070	NM 017638	GK001 NM_020198	HSPC171 NM 014187
BAT2	NM 004638	CDK8	NM 001260	DESC1	NM 014058	FLJ10287	NM 019083	FLJ20080	NM 017657	GLA NM 000169 GLTSCR2 NM 015710	HSPC182 NM 014188 HSPC189 NM 016535
BAT3 BAT4	NM_004639 NM_033177	COKN16 CEBPA	NM_004064 NM_004364	DGUOK DIS3			NM, 018061	FLJ20081 FLJ20084	NM 017658 NM 017659	GNAI3 NM 006496 GNB2L1 NM 006098	HSPE1 NM_002157 HSU79274 NM_013300
BAZIB	NM 032403	CEBPB	NM 005194	DJ37E16.5	NA1 020315	FLJ10374	NM 018074	FLJ20125	NM 017676	GNS NM 002076	HSU84971 NM 013303
BCAR1 BCCIP	NM 014587 NM 016587	CEP2 CES2	NM_006779 NM_003869	DKFZP434B1{ DKFZP434C2;	NM_015426			FLJ20189 FLJ20190	NM 017704 NM 017705	GOSR1 NM_004871 GOSR2 NM_004287	HT010 NM 018471 HT011 NM 018472
BCKDHA BCL2L1	NM 000709	CETN2 CETN3	NM_004344	DKFZp434E22 DKFZP434E23	NM_017612	FLJ10415	NM 018089	FLJ20257	NM_019606	GOT1 NM 002079	HUNK NM_014586
BCS1L	NM 004328	CFL1	NM 005507	DKFZP434L11	NM 032146	FLJ10432	NM 019070	FU20288 FLJ20291	NIM 017748	GPCR150 NM 014373 GPR105 NM 014879	IER5 NM 016545
BET1 BET3	NM 005868 NM 014408	CG005 CG1I	NM_014887 NM_006349	DKFZp434N0£ DKFZp434N14	NM 032261 NM 032133	FLJ10450 FLJ10468	NM_018095 NM_018101	FLJ20342	NM_017774	GFR37 NM 005302	IFRD2 NM 006764 IGBP1 NM 001551
		,				j 1 + 1 44			-mi wiiity	1114 VVVVV4	Tropic Min Min 1991

Fig. 19B

IGSF8 NM_052868 LOC51075 NM_015959 MFAP1 NM_005926 MRPL33 NM 004891 NTE NM 006702 PPP1R158 NM 032	
IMAGE145052 NM_014267 LOC51076 NM 015960 MGC10433 NM 024321 MRPL43 NM 032112 NTF2 NM 005796 PPP2R58 NM 005	
HIAGE345520 NM 024005 LOC51077 NM 015962 MGC10500 NM 031477 MRPL44 NM 022915 NUCB1 NM 006184 PPP6C NM 003	2721 Rpo1-2 NM 019014
IMMT NM 006839 LCC51094 NM 015999 MGC10702 NM 032663 MRPL46 NM 022183 NUDT2 NM 001161 PRCC NM 001101 NM 016053 NM 014652 LCC51096 NM 016001 MGC10924 NM 030571 MRPL48 NM 016055 NUDT5 NM 014142 PRDM5 NM 018	5973 RPS14 NM 005617 8699 RPS18 NM 001020
INCENP NM_020238 LOC51104 NM_016014 MGC10974 NM_032305 MRPL51 NM_016497 NUDT6 NM_007083 PRDX5 NM_012	2094 RPS18 NM, 022551
ING4 NM 016162 LOC51117 NM 016035 MGC11102 NM 032325 MRPS11 NM 022839 NUP54 NM 017426 PRKCABP NM 012	
INVS NM 014425 LOC51118 NM 016037 MGC11115 NM 032310 MRPS12 NM 021107 NUP62 NM 012346 PRKCE NM 0021184 NM 003604 LOC51142 NM 016139 MGC11266 NM 024322 MRPS14 NM 022100 NVL NM 002533 PRO2389 NM 022100 NVL	
TTGA6 NM 000210 LOC51174 NM 016261 MGC1127 NM 033549 MRPS15 NM 031280 NYD-SP11 NM 031851 PRP18 NM 003	3675 RPS27A NM_002954
ITGA9 NM 002207 LOC51187 NM_016304 MGC11279 NM_024326 MRPS18 NM_016065 NY-REN-41 NM 080654 PRPF31 NM_016B3BP NM 014288 LOC51202 NM 016355 MGC11296 NM 032352 MRPS18B NM 014046 OBTP NM 013397 PRRG2 NM 00000000000000000000000000000000000	5629 RPS28 NM_001031 1951 RPS3 NM 001005
ITM1 NM 002219 LOC51204 NM 016360 MGC11352 NM 030927 MRPS18C NM 016067 OGFR NM 007346 PRSS25 NM 013	3247 RPS3A NM_001008
JTB NM_005694 LOC51231 NM_016440 MGC12981 NM_032357 MRPS23 NM_016070 OPA3 NM_025136 PSMA1 NM_002	2786 RPS6 NM_001010
KARS NM 005548 LOC51246 NM 016479 MGC13102 NM 032323 MRPS27 NM 015084 ORC1L NM 004153 PSMA2 NM 007 NB 007 NM 020345 LOC51287 NM 016565 MGC13114 NM 032366 MRPS28 NM 014018 ORC3L NM 012381 PSMA3 NM 007 NM 007 NM 012381 PSMA3 NM 007 NM 017 NM	
KCNQ5 NM 019942 LOC51290 NM 016570 MGC13138 NM 033410 MRPS30 NM 016640 OSBP NM 002556 PSMA5 NM 002	2790 RPS6KC1 NM_012424
KIAA0028 NM_015340 LOC51300 NM_016589 MGC1346 NM_032758 MRPS7 NM_015971 OSCAR NM_130771 PSMB5 NM_002	
KIAA0057 NM 012288 LOC51326 NM 016632 MGC14126 NM 032899 MSMB NM 002443 OSGEP NM 017807 PSMB7 NM 002 KIAA0092 NM 014679 LOC51329 NM 016638 MGC14151 NM 032356 MSTP028 NM 031954 P125 NM 007190 PSMC4 NM 008	2799 RRP46 NM 020158
KIAA0102 NM_014752 LOC51596 NM 015921 MGC14288 NM 032901 INTERF NM 006980 P15-2 NM 018698 PSMD1 NM_002	2807 RXRB NM_021976
KIAA0105 NM_004906 LOC51604 NM_015937 MGC14421 NM_032907 MTF1. NM_005955 P29 NM_015484 P5MD10 NM_002 KIAA0164 NM_014739 LOC51605 NM_015939 MGC14595 NM_032334 MTHFD1 NM_005956 P5326 NM_031450 P5MD4 NM_002	2814 SACM2L NM 022553 2810 SAD1 NM 006590
KIAA0196 NM 014846 LOC51626 NM 016008 MGC14697 NM 032747 MTMR4 NM 004687 PACE NM 002569 PSMD7 NM 002	2811 SAP18 NM 005870
KIAA0255 NM_014742 LOC51631 NM_016019 MGC14798 NM_080650 MTR NM_080254 PACE4 NM_002570 PSMD8 NM_080258 NM_014785 LOC51633 NM_016023 MGC14836 NM_033412 MTRF1 NM_084294 PAFAH2 NM_08437 PSME3 NM_0858 NM_08	
KIAA0274 NM 014845 LOC51644 NM 016057 MGC15677 NM 032878 MITTE NM 016020 PAI-RBP1 NM 015640 PTD009 NM 016 KIAA0317 NM 014821 LOC51651 NM 016077 MGC16169 NM 033115 MUT NM 000255 PANX2 NM 052839 PTD012 NM 014	3146 SBP2 NM 024077
KIAA0372 NM 014639 LOC51657 NM 016086 MGC16386 NM 080668 MUTYH NM 012222 PAPA-1 NM 031288 PTD013 NM 015	5952 SCML1 NM_006748
KIAA0391 NM 014672 LOC51691 NM 016200 MGC16733 NM 033547 MXII NM 005962 PARVB NM 013327 PTD015 NM 014 KIAA0416 NM 015564 LOC54516 NM 019041 MGC17347 NM 138333 MYCBP NM 012333 PAWR NM 002583 PTK7 NM 002	1040 SCYE1 NM 004757 2821 SDCCAG10 NM 005869
KIAA0419 NM. 014711 LOC54543 NM 019059 MGC19595 NM_033415 MYL6 NM 079424 PAX1 NM 005192 PTPN13 NM, 006	264 SDCCAG28 NM_006645
KIAA0433 NM 015216 LOC55954 NM 019103 MGC2404 NM 032360 NAGK NM 017567 PCQAP NM 015889 R3HDM NM 015	
KIAA0438 NM 014819 LOC56851 NM 020154 MGC2408 NM 032331 NAKAP95 NM 014371 PCYT1A NM 005017 RA410 NM 016 KIAA0537 NM 014840 LOC56902 NM 020143 MGC24447 NM 138288 NAPA NM 003827 PDCD10 NM 007217 RAB11A NM 004	106 SDHC NM 003001
KIAA0547 NM 014793 LOC56993 NM 020243 MGC2474 NM 023931 NBP NM 025233 PDE4DIP NM 014644 RAB18 NM 021	252 SEC22L1 NM 004892
KIAA0670 NM 014977 LOC57019 NM 020313 MGC2477 NM 024099 NBR2 NM 005821 PDE6D NM 002601 RAB1B NM 030 NIAA0682 NM 014852 LOC57107 NM 020381 MGC2488 NM 024039 NCBP1 NM 002488 PDE9A NM 002606 RAB2 NM 002	
KIAA0710 NM 014871 LOC57109 NM 020385 MGC2508 NM 024327 NCBP2 NM 007362 PEAS NM 057161 RAB30 NM 014	488 SEC8 NM_021807
KIAA0795 NM 025010 LOC63929 NM 022098 MGC2650 NM_024108 NDUFA1 NM 004541 PEMT NM 007169 RAB7 NM_004	637 SEL1L NM 005065
KIAA0806 NM 014813 LOC81034 NM 030780 MGC2655 NM 024339 NDUFA3 NM 004542 PET112L NM 004564 RABAC1 NM 006 KIAA0872 NM 014940 LOC81558 NM 030802 MGC2747 NM 024104 NDUFA4 NM 002489 PEX11B NM 003846 Rabio4R NM 017	1423 SENP1 NM 014554
KIAA0907 NM 014949 LOC89953 NM 138343 MGC2840 NM 024078 NDUFA5 NM 005000 PEX12 NM 000286 RAD51 NM 138	487 SERPINB3 NM 006919
KIAA0950 NM 012306 LOC90346 NM 138351 MGC3121 NM 024031 NDUFA6 NM 002490 PEX13 NM 002618 RAGA NM 008 KIAA0971 NM 014929 LOC90678 NM 138361 MGC3123 NM 024107 NDUFA7 NM 005001 PEX16 NM 057174 RAI2 NM 021	570 SERPINB8 NM_002640 785 SERPINI1 NM 005025
[KIAA1012 NM, 014939 [LOC90701 NM, 033280 [MGC3133 NM, 031287 [NDUFB3 NM, 002491] PEXB NM, 000287 [RAMP NM, 016	448 SES2 NM_031459
KIAA1041 NM 014947 LOC92105 NM 138381 MGC3222 NM 024334 NDUFS1 NM 005006 PHACS NM 032592 RANGAP1 NM 002	1883 SF3A3 NM_006802
KIAA1068 NM 015332 LRP5 NM 002335 MGC3248 NM 032486 NDUFS3 NM 004551 PHB NM 002634 RAP1 NM 018 KIAA1100 NM 014901 LRRN1 NM 002319 MGC4054 NM 024341 NDUFS4 NM 002495 PHKB NM 000293 RARG-1 NM 016	975 SF3B1 NM 012433 167 SF3B2 NM 006842
[KIAA1608 NM_024820 LSM3 NM 014463 MGC4093 NM_030578 NDUFV1 NM 007103 PIGN NM 012327 RASSF1 NM 007	182 SF3B4 NM 005850
KIF3B NM_004798 LSM5 NM_012322 MGC4189 NM_032308 NEK7 NM_133494 PIGPC1 NM_022121 RBBP4 NM_005	
KIF9 NM_022342 LTA4H NM_000895 MGC4251 NM_032376 NFATC2 NM_012340 PIGPC1 NM_022121 RBL1 NM_002 KLRF1 NM_016523 LYPLA2 NM_007260 MGC4308 NM_032359 NFE2L1 NM_003204 PIK3C3 NM_002647 RBL2 NM_005	895 SFRS2 NM_003016
KNSL7 NM 020242 LZTFL1 NM 020347 MGC4608 NM 024516 NFE2L3 NM 004289 PINK1 NM 032409 RBM15 NM 022	768 SFRS8 NM 004592
KPTN NM 007059 LZTR1 NM 006767 MGC4767 NM 032314 NFKBIB NM 002503 PIP5K1A NM 003557 RBM6 NM 005 KRT10 NM 000421 M17S2 NM 031858 MGC4771 NM 032668 NFKBIB NM 002503 PIST NM 020399 RBM7 NM 016	777 SGCE NM_003919 090 SGT1 NM_006704
LAPTM4A NM 014713 M6A NM 019852 MGC5302 NM 024089 NFKBIB NM 002503 PL6 NM 007024 RDBP NM 002	904 SH3BGRL2 NM 031469
LCP NM_014315 M96 NM 007358 MGC5378 NM_032632 NFKBIL1 NM_005007 PLA2G4B NM_005090 REA NM_007	273 SIP NM, 014412
LDB1 NM_003893 MAGOH NM_002370 MGC5469 NM_032361 NFYA NM_002505 PLAA NM_004253 RECQL NM_002 LEPR NM_002303 MAP2K5 NM_002757 MGC5609 NM_024093 NIMP NM_032730 PLDN NM_012388 RECQL5 NM_004	
LGMN NM 005606 MAP3K11 NM 002419 MGC5521 NM 024061 NKTR NM 005385 PME-1 NM 016147 REG1A NM 002	909 SKD1 NM 004869
LHX6 NM_014368 MAP3K3 NM_002401 MGC9084 NM_033418 NLN NM_020728 PMS2 NM_000535 REG1B NM_006 LIM NM_006457 MAP3K7 NM_003188 MGC9740 NM_080658 NMA NM_012342 PMS2LB NM_005394 RENT1 NM_002	911 SKI NM_003036
LIMS1 NM 004987 MAPK7 NM 002749 MGST3 NM 004528 NME1 NM 000269 PNAS-131 NM 031446 RFC3 NM 002 LIN-7-C NM 018362 MAPK8IP2 NM 012324 MID1 NM 000381 NME7 NM 013330 PNKP NM 007254 RFPL2 NM 005	915 SKP2 NM 032637
LISCH7 NM 015925 MAPK8IP3 NM 015133 MKRN1 NM 013446 NOH61 NM 019082 PNMA1 NM 006029 RNF40 NM 014	771 SLC25A19 NM 021734
LIV-1 NM 012319 MAT2A NM 005911 MLH1 NM 000249 NOLA1 NM 018983 PODXL NM 005397 RNF5 NM 006 LOC113251 NM 052879 MBD4 NM 003925 MLN NM 002418 NOLC1 NM 004741 POLE3 NM 017443 RNGTT NM 003	
LOC113444 NM 138428 MCEE NM 032601 MN1 NM 002430 NOSIP NM 015953 POLL NM 013274 RNPC2 NM 004 LOC113622 NM 138430 MCFP NM 018843 MOCS3 NM 014484 NOT56L NM 005787 POLR2A NM 000937 RPA2 NM 002	902 SLC35A1 NM_006418
LOC115827 NM 138453 MCM3 NM 002388 MPPE1 NM 023075 NPAS2 NM 002518 POLR2K NM 005034 RPA40 NM 004	875 SLC7A9 NM 014270
LOC129401 NM 13B285 MDFI	
LOC153768 NM_138492 MDH2 NM_005918 MRPL18 NM_014161 NR1D1 NM_021724 POP5 NM_015918 RPL18 NM_000	979 SMARCA5 NM 003601
LOC51002 NM 016058 MDS025 NM 021825 MRPL19 NM 014763 NR1H3 NM 005693 POR1 NM 0124D2 RPL18A NM 000 LOC51004 NM 015940 MDS032 NM 018467 MRPL2 NM 015950 NRAS NM 002524 POU5F1 NM 002701 RPL28 NM 000	987 SMC1L1 NM_008306
LOC51016 NM 016049 MDS033 NM 018468 MRPL22 NM 014180 NRCAN NM 005010 PPIL1 NM 016059 RPL27 NM 000	988 SMC2L1 NM_008444
LOC51026 HM_016072 MEN1 NM_130800 MRPL27 NM_016504 NS1-BP NM_006469 PPP1CA NM_002708 RPL32 NM_000	994 SMCX NM_004187
LOC51027 NM 016074 MEP50 NM 024102 MRPL3 NM 007208 NSEP1 NM 004559 PPP1R10 NM 002714 RPL37 NM 000 LOC51060 NM 015913 METAP2 NM 006838 MRPL30 NM 016503 NSF NM 008178 PPP1R11 NM 021959 RPL37A NM 000	997 SMPD2 NM 003080
LOCS1067 NM 015936 METL NM 018396 MRPL32 NM 031903 NT5C3 NM 016489 PPP 1R12B NM 032105 RPL41 NM 021	104 SNRPD2 NM 004597

Fig. 19C

Gene Name ReiSeq Y Gene SNRPD3 NM 004175 TXNL SNRPF NM 003095 U2AF1 SNW1 NM 012245 U5-100 SNX1 NM 003099 U5-110 SNX11 NM 013323 UBE20 SNX17 NM 014748 UBE20 SNX5 NM 014426 UBE20 SON NM 003103 UBO	Name : Proportion
SNRPF NM 003095 U2AF	DBCIBN ATAL, COMPAN
CARLLA ALCA ALCA I	NM 0047 NM 0067
SNW1 NM_012245 U5-100	X NM_0048
SNX1 NM_003099 U5-11	KD NM_0042
SNX11 NM 013323 UBE21	4 NM 0039
SNX17 NM_014748 UBE28 SNX5 NM_014426 UBE28	NM_003:
SNX5 NM 014426 UBE21 SON NM 003103 UBQL	/1 NM_0224 N1 NM_053(7 NM_0159 EL1 NM_0058
SOX17 NM 022454 UCH3	V1 NM 053(7 NM 0159
SOX9 NM 000346 UGTR	EL1 NM_0058
SP2 NM_138406 UMPS	NM_0003
SPATA2 NM 006038 UNRIP	NM_0071
SPC18 NM 014300 UPF3E SPG4 NM 014946 UQCR	NM 0806
SPG4 NM_014946 UQCR SPK NM_004819 UQCR	C2 NM_0035
SORDL NM 021199 URKL1	H NM_006C NM_0178
SRP19 NM 003135 UROD	NM_0003
SRP54 NM_003136 UROS	NM 0003
SRP68 NM_014230 USF1	NM .0071
SSA2 NM_004600 USP5 SSBP1 NM_003143 UXT	NM 0034
SSBP1 NM_003143 UXT SSFA2 NM_006751 V1RL1	NM 0041
\$\$R2 NM_003145 VEGFO	, NM 0206 NM, 005
SSR3 NM_007107 VMP1	NM_030\$
SSSCA1 NM_006396 VPS33	NM_030\$ NM_022\$
SSTK NM 032037 WARS	NM 0158
SSTR4 NM_001052 WBP4 ST13 NM_003932 WDF2	NM 0071
ST13 NM_003932 WDF2 STAF42 NM_053053 WDR12	NM 0529
STAF65(gami NM 014860 WDR13	NM_0182 NM_017E
STAM NM 003473 WHIP	NM 0201
STAM2 NM.005843 XPC STCH NM.006948 XPO1	NM_0046
STCH	NM 0034
STK19 NM_004197 XRCC4 STK24 NM_003576 XRCC5	NM_0228
STOML1 NM_004809 XRN2	NM_0211 NM_0122
STOML2 NM 013442 YR-29	NM 0148
STX18 NM 016930 YWHAE	NM 0034
SUCLG1 NM_003849 ZBRK1	
SULT1A3 NM 003166 ZF5128 SULT1C1 NM_001056 ZFP37	
SULT1C1	NM 0034
SUPV3L1 NM 003171 ZFP95	NM_0042 NM_0145
T54 NM 015698 ZNF133	
TADA3L NM_133480 ZNF134	NM_0034
TAF11 NM 005643 ZNF142	NM_005(
TAF6 NM 005641 ZNF146 TARBP2 NM 004178 ZNF155	
TARBP2 NM_004178 ZNF155 TAX18P1 NM_006024 ZNF175	
TCERG1 NM 006708 ZNF183	
TCF1 NM_000545 ZNF189	
TCF2 NM_000458 ZNF192	NM_0062
TCF2 NM_000458 ZNF193	
TCF2 NM 000458 ZNF207 TCOF1 NM 000356 ZNF214	
TCP1 NM 030752 ZNF221	
TDRKH NM 006862 ZNF222	
TEGT NM_003217 ZNF224	NM 0133
TESK2 NM_007170 ZNF225	
TFAP4 NM 003223 ZNF226	
TFPT NM 013342 ZNF230 TG737 NM 006531 ZNF264	
TIMM23 NM 006327 ZNF265	
TIMM9 NM 012460 ZNF277	NM 0218
TIP39 NM 012143 ZNF300	NM .0528
TLE3 NM_005078 ZNF302 TLN1 NM_006289 ZNF304	
TLN1 NM_006289 ZNF304 TM9SF1 NM_006405 ZNF317	
TM9SF2 NM_004800 ZNF338	
TMOD2 NM_014548 ZNF345	
TMP21 NM 006827 ZNF361	NM 0185
TMSB10 NM 021103 ZNF-U69	
TNFAIP1 NM_021137 ZNRD1 TOMM70A NM_014820	NM ,0145
TOR2A NM 130459	,
TPT NM_014317	
TRA1 NM_003299	
TRA1 NM 003299 TRAF5 NM 004619	
TRA1 NM 003299 TRAF5 NM 004619 TRAP150 NM 005119	
TRA1 NM 003299 TRAF5 NM 004619 TRAP150 NM 005119 TRFP NM 004275	
TRA1 NM 003299 TRAF5 NM 004619 TRAP150 NM 005119 TRFP NM 004275 TRIM4 NM 033017 TRIP NM 005879	
TRA1 NM_003299 TRAF5 NM_004619 TRAP150 NM_005119 TRFP NM_004275 TRIM4 NM_033017 TRIP NM_005879 TRIP11 NM_004239	
TRA1 NM_003299 TRAF5 NM 004619 TRAP150 NM 005119 TRFP NM 004275 TRIM4 NM 033017 TRIP NM 005879 TRIP11 NM_004239 TRN-SR NM_012470	
TRA1 NM_003299 TRAF5 NM 004619 TRAP150 NM 005119 TRFP NM 004275 TRIM4 NM 033017 TRIP NM 005879 TRIP11 NM_004239 TRN-SR NM_012470 TRPS1 NM 014112	
TRA1 NM 003299 TRAF5 NM 004619 TRAP150 NM 005119 TRFP NM 004275 TRIM4 NM 033017 TRIP NM 005879 TRIP11 NM 004239 TRN-SR NM 012470 TRPS1 NM 014112 TSG101 NM 006292	
TRA1 NM_003299 TRAF5 NM 004619 TRAP150 NM 005119 TRFP NM 004275 TRIM4 NM 033017 TRIP NM 005879 TRIP11 NM_004239 TRN-SR NM_012470 TRPS1 NM 014112	
TRA1 NM 003299 TRAF5 NM 004619 TRAP150 NM 005119 TRFP NM 004275 TRIM4 NM 033017 TRIP NM 005879 TRIP11 NM 004239 TRN-SR NM 012470 TRPS1 NM 014112 TSG101 NM 006292 TSLRP NM 012472	

(32/41)

Fig. 20A

· [:			2 2						
. 1		C-2-	• •	1		HNE1a	/HNF4a-	•	,•
		URKI.1	NM_017859	SCYE1	NM_004757	RPL37AP1	NG_000988	FE65L2	NM_006051
		FLJ20671		C1S		XGH	NM_000613	M17S2	NM_031858
	NM_025115	FLJ21963	NM_024560	G3A		FL J 10278	NM_018045	FLJ10583	NM_018148
		HNT48/	Ar50946/	MGC15435		ACP3	NM_004925	AIFZ	NM_001880
	197770 WW	LUC5128/		MGC11034		NGKZ XDI.	072910 MN	EST CST	NM 013324
	NM_UZUSBB	PHTE	NW_014546	CF52	NM_010413	AUT El 14000	MM_0005/8	GPXZ CEDDIMA10	NM_002083
	NM 025192	RPS6KAS		F1.112788		AGT	NM_018233	SECTIVATO HOCECO	NW_016166
	NM 001080	TM4SF4		LOC56902		APOA2	NM 001643	HNMT	NM 006895
		GRO3		HSPC111		GOS2	NM 015714	SI C17A2	NM 005835
	NM_002568	UGT2B15		NR5A2		WBP4	NM 007187	APCS	NM 001639
	NM_001798	FXYD7	NM_022006	FLJ13611	NM 024941	ELF3	NM 004433	NFKBIB	NM 002503
	NM_000392	FLJ12770		ABCC2		PAFAH2	NM 000437	FLJ11838	NM 024664
	NM_000043	FU22169	NM_024085	TNFRSF6	NM_000043	SSTR1	NM_001049	HPCL2	NM_012260
	NM_001073	FLJ10415		UGT2811	NM_001073	PIST	NM_020399	WDR12	NM_018256
	NM_017909	CYP3A43	NM_022820	C4BPA	NM_000715	PLGL	NM_002665	LOC51096	NM_016001
		ACVR1		GTF2E1	NM_005513	C8B	990000 WN	SERPINE	NM_000602
	NM_000063	SNW1		BAT3	NM_004639	MGC11266	NM_024322	(MT1X	NM_005952
		REA	NM_007273	23		LOC51326	NM_016632	CLYBL	NM_138280
		M17S2		ADH6	NM_000672	1,000,000	NM_015913	CYB5-M	NM_030579
	NM_017659	၁၉၁		FLJ20080	NM_017657	FLJ13448	NM_025147	MTHFD1	NM_005956
IKF6 DKE2056400522	NM_U0514/	NOLC:	NM_004/41	APG3	NM_022488	ASGR1	NM_001671	SSA2	NM_004600
3		APCS		MRPS188	NA 010204	462	NIM 003613	HUGV	NW DOODS
	NM_032982	WDR12		LOC54518		ZNF361	NM 018555	FL 122551	NM 024708
	NM_000062	FACTP140	NM_007192	AP3M1	NM_012095	CTSZ	NM_001336	T1H4	NM_002218
ADH18	NM_000668	KIAA0806	•	AMBP		PAX8	NM_013952	NRD1	NM_002525
	ZOSOSO MIN	OAZ F	NM_00233/	SEL1L	NM_005065	K!AA1041	NM_014947	SAC SAC SAC SAC SAC SAC SAC SAC SAC SAC	NM_018417
	NM_UUDS22	מן מסק	017100_MN	HSPC129	NM_016396	IIIH3	VM_002217	I AKS	NM_003191
	NM 000481	STON CITAN		SERBING4	NIM COUNTRY	FMAD1	NM 000629	75.0	NM 0001/3
	NM 002040	TAT		ADH18	NM OODSER	HNF427	AF509467	100 1100	NM 003210
	NW 005800	CNS		MEDI 45	NN 014175	ANDED	MM 001150	Coco	MM_002079
		Spi P1		HADI	NM 017545	IGERP1	NM 000596	ASGP2	NW_001310
DKFZP586A0522	NM 014033	S281		SYN3	NM 133632	RAMP		Self.	NM 000166
	NM_005826	HBP1	NM_012257	AHSG	NM_001622	SERPINA1	NM_000295	RBP5	NM 031491
	NM_004766	CLDN2	NM_020384	MTP	NM_000253	SUPV3L1	NM_003171	STRAIT11489	NM_021242
		AASS	NM_005763	AUTL1	NM_032852	五		INADL	NM_005799
	NM_016023	PSMA1	NM_002786	RAB33B	NM_031296	TM4SF4		MT1L	
	NM_003/42	ALDH3A1		DAF		NAPA		M96	NM_007358
	NIM_001818	11.147.5	NM_002651	PCK1	NM_002591	SEKPINCI	NM_U000488	FU21272	NM 025032
	NM 024743	SERPINI	NM 005025	NROP2	N.M. 021969	HAI		1 282	NA 000567
		<u>E</u>		НСО		NET:2	NM 012338	SERPINAG	NM 001756
	1	CRP		DUSP6	NM_022652	MT1H		ACF	NM_014576
		EIF4E	NM_001968	GBE1	NM_000158	CYBS	NM_001914	DKFZP56400463	NM_014156
	NM_007114	GSS	NM_000178	AKR1C2	NM_001354	DC13	NM_020188	HSD1782	NM_002153
	NM_004528	FLJ12910	NM_024573	ARHGAP11A	NM_014783	ALS2	NM 020919	616615	מומו חח ומואנ
	NM_015638	FLJ10407	NM_018087	FU10525	NM_018126	RNGTT	NM_003800		
	NM_024773 NM_016281	TAF2GL Sil V	NG_001012 NM_006928	FL)10774 AKR 104	NM_024662 NM_001818	CYP3A43	NM_022820 NM_003064		
		CRADO	NM 003805	PZP		HSD11B1			

Fig. 20B

Feedforward Loop

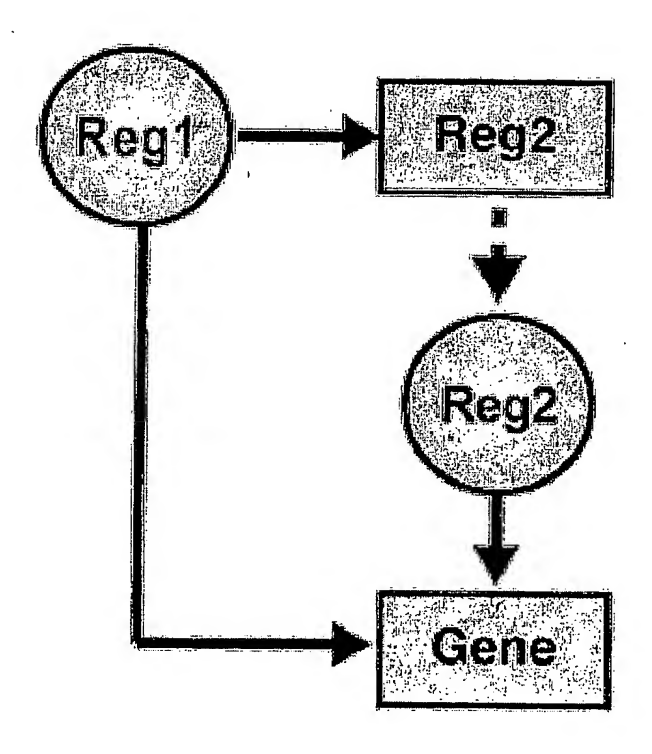


Fig. 21A

Reg1	HNF	-6] H	INF6
Reg2	HNF	4α	H	NF1α
Reg3	HNF	1α		
	C1S	NM_001734	F11	NM_019559
. : •	ABCC2	NM_000392	C1S	NM_001734
.3 () () ()	TNFRSF6	NM_000043	FLJ10650	NM_018168
	UGT2B11	NM_001073	ABCC2	NM_000392
	C2	NM_000063	TNFRSF6	NM_000043
	AMBP	NM_001633	UGT2B11	NM_001073
	SERPING1	NM_000062	UGT1A1	NM_000463
	ADH1B	NM_000668	C2	NM_000063
	PCK1	NM_002591	ADH1A	NM_000667
•	DKFZP586A0522	NM_014033	AMBP	NM_001633
	VTN	NM_000638	SERPING1	NM_000062
S	AKR1C4	NM_001818	ADH1B	NM_000668
	FLJ21934	NM_024743	HABP2	NM_004132
moters	KIAA0872	NM_014940	PCK1	NM_002591
) 	RPL37AP1	NG_000988		NM_014033
	PLGL	NM_002665	VTN	NM_000638
2	C8B	NM_000066	AKR1C4	NM_001818
	LOC51060	NM_015913	FLJ21934	NM_024743
und	HNF4a7	AF509467	KIAA0872	NM_014940
\subseteq	TM4SF4	NM_004617	RPL37AP1	NG_000988
0	UGT2B15	NM_001076	PLGL	NM_002665
m	CYP3A43	NM_022820	C8B	NM_000066
	M17S2	NM_031858	LOC51060	NM_015913
	HNMT	NM_006895	HNF4a7	AF509467
	APCS	NM_001639	TM4SF4	NM_004617
·	WDR12	NM_018256	UGT2B15	NM_001076
	APOH	NM_000042	CYP3A43	NM_022820
	GJB1	NM_000166	M17S2	NM_031858
·	CRP	NM_000567	HNMT	NM_006895
			APCS	NM_001639
			WDR12	NM_018256
			APOH	NM_000042
				NM_000166
i		ļ	CRP	NM 000567

Fig. 21B

Multi-input

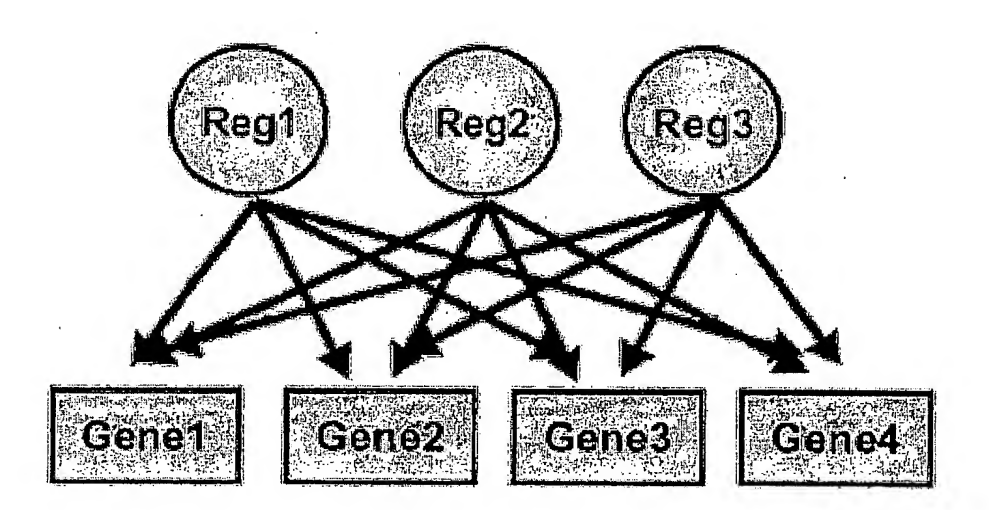


Fig. 22A

Reg1		H	NF6		HNF1α	/HNF4α
Reg2	•	HI	NF4α	1,	HNF4α	/HNF1a
2553年3143	BCKDHA	NM_000709	FLJ13798	NM_024773	FLJ13273	NM_024751
	FLJ23263	NM_025115	GSS	NM_000178	MGC10500	NM_024701
	FLJ11271	NM_018373	НВОА	NM_007067	SDCCAG10	NM_005869
<u> </u>	HMG2	NM_002129	LOC51060	NM_015913	FBXO8	NM_012180
	LOC81558	NM_030802	FLJ13220	NM_021927	ZNF300	NM_052860
	SAS10	NM_020368	FLJ12910	NM_024573	H4F2	NM_003548
	SEC10L1	NM_006544	FLJ10407	NM_018087	FLJ11301	NM_018385
	RRP46	NM_020158	FLJ10342	NM_018064	SEL1L	NM_005065
'	SNRPD2	NM_004597	FLJ20671	NM_017924	ZNF155	NM_003445
	MDH1	NM_005917	LOC51287	NM_016565	C6orf11	NM_005452
	ORC1L	NM_004153	GLA	NM_000169	ARHGAP11A	NM_014783
	FLJ20627	NM_017909	RPS6KA5	NM_004755	UROD	NM_000374
	GTF2E1	NM_005513	FLJ20772	NM_017956	FLJ20731	NM_017946
	TOMM70A	NM_014820	FLJ12770	NM_032174	RAB6KIFL	NM_005733
	PAPA-1	NM_031288	FLJ22169	NM_024085	TMP21	NM_006827
်လ	HASJ4442	NM_017528	FLJ10415	NM_018089	MGC15677	NM_032878
<u>a</u>	FLJ20084	NM_017659	ZNF317	NM_020933	WBP4	NM_007187
romote	PEX6	NM_000287	SNW1	NM_012245	PAFAH2	NM_000437
Ε	FLJ11301	NM_018385	REA	NM_007273	EIF3S6	NM_001568
9	EED	NM_003797	C2F	NM_006331	PSMA5	NM_002790
ā	MGC19595	NM_033415	NOLC1	NM_004741	TMOD2	NM_014548
O	CIR	NM_004882	CLONE24922	NM_015679	GLA	NM_000169
l l	CLLD8	NM_031915	ССТ8	NM_006585	GNB2L1	NM_006098
O	ABCB8	NM_007188	PSMB1	NM_002793	FNTB	NM_002028
Bound	SPG4	NM_014946	WDR12	NM_018256	PEX13	NM_002618
	GA8PA	NM_002040	KIAA0806	NM_014813	FE65L2	NM_006051
	OGFR	NM_007346	DKFZp761J139	NM_032280	UQCRC2	NM_003366
• •	COPB2	NM_004766	SART3	NM_014706	FLJ14855	NM_033210
	AF15Q14	NM_020380	COX7A2L	NM_004718	HHLA2	NM_007072
	MTERF	NM_006980	FLJ20422	NM_017814	CYB5-M	NM_030579
	LOC51633	NM_016023	COPS7A	NM_016319	CDC45L	NM_003504
	FLJ14486	NM_032792	FLJ20643	NM_017916	panp	NM_020357
	FLJ21934	NM_024743	HBP1	_	FLJ20643	NM_017916
. · · ·	KIAA0872	NM_014940	PSMA1		FLJ21272	NM_025032
	TEGT	NM_003217	FLJ21272	NM_025032		
	MGC4189	NM_032308	FLJ11029	NM_018304		
	SERPINB8	NM_002640	ARL1	NM_001177		
	MGST3	NM_004528	SERPINI1	NM_005025		
	HSP1058	NM_006644	NUDT2	NM_001161		
	C20orf188	NM_015638				

Table S11. The feed forward regulatory motifs in pancreatic islets. The regulatory modules here were derived as described in Supporting Online Material. Feed forwards only involving HNF1 α and HNF4 α are also multi-input motifs, as they bind each other's promoters in a multicomponent loop.

Fig. 22B

Feedforward Loop

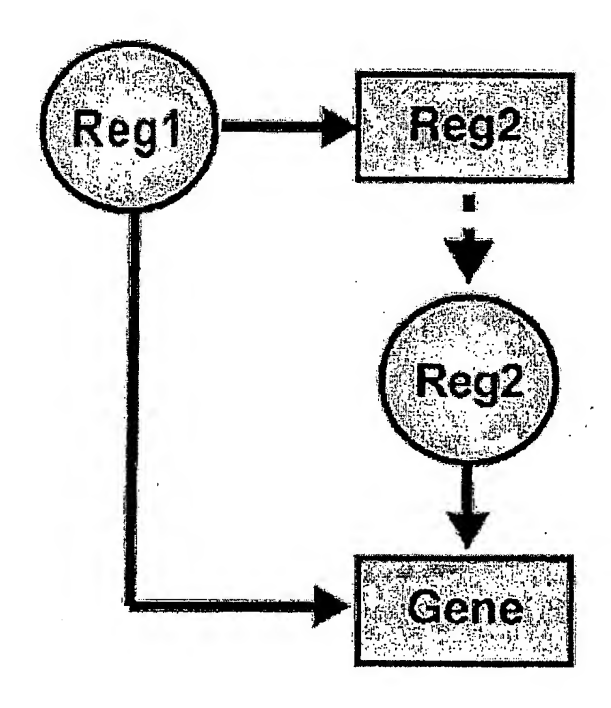


Fig. 23A

INF10 HNF6 HR740	01 NM_018385	NM_000169	43 NW_017916	72 NM_025032							
	FLJ11301	GLA	FLJ20643	FLJ21272							
TNT C	NM_018168	NM_020147	NM_018385	NM_021969	NM_030967	AF509467	NM_017691	NM_000169	NM_000042	NM_017916	NM 025032
	FLJ10650	LOC56906	FLJ11301	NR0B2	KRTAP1.1	HNF4a7	FLJ20156	GLA	APOH	FLJ20643	FLJ21272
		S	l O]	Ol	UC)ار	J [)U	nc	В	
Reg1 Reg2 Reg3	1		Maria in res	· .							

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Fig. 23B

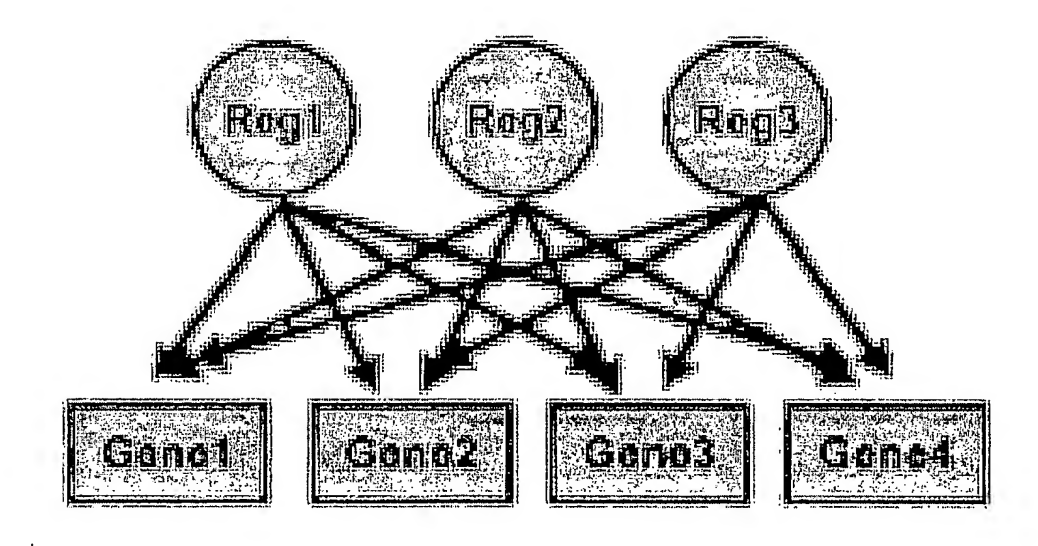


Fig. 24

		T	Hepatocytes	ytes			Pancr	Pancreatic Islets	Islets
		HINIER					HNFA		HNF10
	¥		1	•	77		•		•
	HNF1A	SP2	NR0B2	TEF	HNF4A	HNF1A	SP2	BLZF1	HNF4A
	HNF1B	NR112	NR5A2	RAMP	NR1D1	HNF1B	CREBL2 MEF2B	MEF2B	ELF3
	LISCH7	SREBF2	CREBL2 ATF2	ATF2		LISCH7	NR1D1	MTF1	PAX8
	RXRB	ВТЕЗ	ELF3	M96		RXRB	LZTR1	CRSP3	NR5A2
Transcription	NR1H3	HIF1A	PAX8			NR1H3	E2F4	HCNGP	NR0B2
Factors	DED	NR3C2				DED	E2F5	NR1H3	NR2C2
	GABPA	TCF19				GABPA	M96	POU5F1	
	GABPB2			•		GABPB2	TFAP4	RAMP	
	ATF4					ATF4	ATF6	USF1	
	ATF7					ATF7	LZTFL1		
	TRAP150 CNOT2	CNOT2				TRAP150	TRIP11	NCOA4	
	TADA3L	CRSP9				FACTP140 CIR	CIR	SMAP	
COACIIVAIOIS						SMARCA5 CNOT3	CNOT3		
						COASTER	CNOT4		
Mitochondrial	mtTFB	TFAM				mtTFB	mtERF		

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